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(54) Title: GLYPHOSATE TOLERANT 5-ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE SYNTHASES

(57) Abstract

Genes encoding class II EPSPS enzymes are disclosed. The genes are useful in producing transformed bacteria and plants which are tolerant to glyphosate herbicide. Class II EPSPS genes share very little homology with known, Class I EPSPS genes, and do not hybridize to probes from Class I EPSPS's. The Class II EPSPS enzymes are characterized by being more kinetically efficient than Class I EPSPS's in the presence of glyphosate. Plants transformed with Class II EPSPS genes are also disclosed as well as a method for selectively controlling weeds in a planted crop field.

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GLYPHOSATE TOLERANT 5-ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE SYNTHASES

This is a continuation-in-part of a copending U.S. patent application having serial number 07/576,537, filed August 31, 1990 and entitled "Glyphosate Tolerant 5-Enolpyruvylshikimate-3-Phosphate Synthases."

BACKGROUND OF THE INVENTION

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This invention relates in general to plant molecular biology and, more particularly, to a new class of glyphosate tolerant 5-enolpyruvylshikimate-3-phosphate synthases.

Recent advances in genetic engineering have provided the requisite tools to transform plants to contain foreign genes. It is now possible to produce plants which have unique characteristics of agronomic importance. Certainly, one such advantageous trait is more cost effective, environmentally compatible weed control via herbicide tolerance. Herbicide-tolerant plants may reduce the need for tillage to control weeds thereby effectively reducing soil erosion.

One herbicide which is the subject of much investigation in this regard is N-phosphonomethylglycine commonly referred to as glyphosate. Glyphosate inhibits the shikimic acid pathway which leads to the biosynthesis of aromatic compounds including amino acids, plant hormones and vitamins. Specifically, glyphosate curbs the conversion of phosphoenolpyruvic acid (PEP) and 3-phosphoshikimic acid to 5-enolpyruvyl-3-phosphoshikimic acid by inhibiting the enzyme 5-enolpyruvylshikimate-3-phosphate synthase (hereinafter referred to as EPSP synthase or EPSPS).

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It has been shown that glyphosate tolerant plants can be produced by inserting into the genome of the plant the capacity to produce a higher level of EPSP synthase in the chloroplast of the cell (Shah et al., 1986) which enzyme is preferably glyphosate tolerant (Kishore et al. 1988). Variants of the wild-type EPSPS enzyme have been isolated which are glyphosate tolerant as a result of alterations in the EPSPS amino acid coding sequence (Kishore and Shah, 1988; Schulz et al., 1984; Sost et al., 1984; Kishore et al., 1986). These variants typically have a higher K_i for glyphosate than the wild-type EPSPS enzyme which confers the glyphosate tolerant phenotype, but these variants are also characterized by a high K_m for PEP which makes the enzyme kinetically less efficient (Kishore and Shah, 1988; Sost et al., 1984; Schulz et al., 1984; Kishore et al., 1986); Sost and Amrhein, 1990). For example, the apparent K_m for PEP and the apparent K_i for glyphosate for the native EPSPS from E. coli are 10 μ M and 0.5 μ M while for a glyphosate tolerant isolate having a single amino acid substitution of an alanine for the glycine at position 96 these values are 220 µM and 4.0 mM, respectively. A number of glyphosate tolerant plant variant EPSPS genes have been constructed by mutagenesis. Again, the glyphosate tolerant EPSPS was impaired due to an increase in the K_m for PEP and a slight reduction of the V_{max} of the native plant enzyme (Kishore and Shah, 1988) thereby lowering the catalytic efficiency (V_{max}/K_m) of the enzyme. Since the kinetic constants of the variant enzymes are impaired with respect to PEP, it has been proposed that high levels of overproduction of the variant enzyme, 40-80 fold, would be required to maintain normal catalytic activity in plants in the presence of glyphosate (Kishore et al., 1988).

While such variant EPSP synthases have proved useful in obtaining transgenic plants tolerant to glyphosate, it would be increasingly beneficial to obtain an EPSP synthase that is highly glyphosate tolerant while still kinetically efficient such that the amount of the glyphosate tolerant EPSPS needed to be produced to maintain normal catalytic activity in the plant is reduced or that improved tolerance be obtained with the same expression level.

Previous studies have shown that EPSPS enzymes from different sources vary widely with respect to their degree of sensitivity to inhibition by glyphosate. A study of plant and bacterial EPSPS enzyme activity as a function of glyphosate concentration showed that there was a very wide range in the degree of sensitivity to glyphosate. The degree of sensitivity showed no correlation with any genus or species tested (Schulz et al., 1985). Insensitivity to glyphosate inhibition of the activity of the EPSPS from the Pseudomonas sp. PG2982 has also been reported but with no details of the studies (Fitzgibbon, 1988). In general, while such natural tolerance has been reported, there is no report suggesting the kinetic superiority of the naturally occurring bacterial glyphosate tolerant EPSPS enzymes over those of mutated EPSPS enzymes nor have any of the genes been characterized. Similarly, there are no reports on the expression of naturally glyphosate tolerant EPSPS enzymes in plants to confer glyphosate tolerance.

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SUMMARY OF THE INVENTION

A DNA molecule comprising DNA encoding a kinetically efficient, glyphosate tolerant EPSP synthase is presented. The EPSP synthases of the present invention reduce the amount of overproduction of the EPSPS enzyme in a transgenic

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plant necessary for the enzyme to maintain catalytic activity while still conferring glyphosate tolerance. This and other EPSP synthases described herein represent a new class of EPSPS enzymes, referred to hereinafter as Class II EPSPS enzymes. Class II EPSPS enzymes share little homology to known bacterial or plant EPSPS enzymes and exhibit tolerance to glyphosate while maintaining suitable K_m (PEP) ranges. Suitable ranges of K_m (PEP) for EPSPS for enzymes of the present invention are between 1-150 μ M, with a more preferred range of between 1-35 μ M, and a most preferred range between 2-25 µM. These kinetic constants are determined under the assay conditions specified hereinafter. The V_{max} of the enzyme should preferably be at least 15% of the uninhibited plant enzyme and more preferably greater than 25%. An EPSPS of the present invention preferably has a Ki for glyphosate range of between 25-10000 μM . The K_i/K_m ratio should be between 3-500, and more preferably between 6-250. The $V_{\rm max}$ should preferably be in the range of 2-100 units/mg (µmoles/minute.mg at 25°C) and the K_m for shikimate-3-phosphate should preferably be in the range of 0.1 to 50 μ M.

Genes coding for Class II EPSPS enzymes have been isolated from three (3) different bacteria: Agrobacterium tumefaciens sp. strain CP4, Achromobacter sp. strain LBAA, and Pseudomonas sp. strain PG2982. The LBAA and PG2982 Class II EPSPS genes have been determined to be identical and the proteins encoded by these two genes are very similar to the CP4 protein and share approximately 84% amino acid identity with it. Class II EPSPS enzymes can be readily distinguished from Class I EPSPS's by their inability to react with polyclonal antibodies prepared from Class I EPSPS enzymes under conditions where other Class I EPSPS enzymes would readily react with the Class I antibodies.

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Other Class II EPSPS enzymes can be readily isolated and identified by utilizing a nucleic acid probe from one of the Class II EPSPS genes disclosed herein using standard hybridization techniques. Such a probe from the CP4 strain has been prepared and utilized to isolate the Class II EPSPS genes from strains LBAA and PG2982. These genes may also be adapted for enhanced expression in plants by known methodology. Such a probe has also been used to identify homologous genes in bacteria isolated *de novo* from soil.

The Class II EPSPS enzymes are preferably fused to a chloroplast transit peptide (CTP) to target the protein to the chloroplasts of the plant into which it may be introduced. Chimeric genes encoding this CTP-Class II EPSPS fusion protein may be prepared with an appropriate promoter and 3' polyadenylation site for introduction into a desired plant by standard methods.

Therefore, in one aspect, the present invention provides a new class of EPSP synthases that exhibit a low K_m for phosphoenolpyruvate (PEP), a high V_{max}/K_m ratio, and a high K_i for glyphosate such that when introduced into a plant, the plant is made glyphosate tolerant such that the catalytic activity of the enzyme and plant metabolism are maintained in a substantially normal state. For purposes of this discussion, a highly efficient EPSPS refers to its efficiency in the presence of glyphosate.

In another aspect of the present invention, a double-stranded DNA molecule comprising DNA encoding a Class II EPSPS enzyme is disclosed. A Class II EPSPS enzyme DNA sequence is disclosed from three sources: Agrobacterium sp. strain designated CP4, Achromobacter sp. strain LBAA and Pseudomonas sp. strain PG2982.

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In a further aspect of the present invention, a nucleic acid probe from an EPSPS Class II gene is presented that is suitable for use in screening for Class II EPSPS genes in other sources by assaying for the ability of a DNA sequence from the other source to hybridize to the probe.

In yet another aspect of the present invention, transgenic plants and transformed plant cells are disclosed that are made glyphosate tolerant by the introduction of a Class II EPSPS gene into the plant's genome.

In a still further aspect of the invention, a recombinant, double-stranded DNA molecule comprising in sequence:

- a) a promoter which functions in plant cells to cause the production of an RNA sequence;
- a structural DNA sequence that causes the production of an RNA sequence which encodes a Class II EPSPS enzyme; and
- c) a 3' nontranslated region which functions in plant cells to cause the addition of a stretch of polyadenyl nucleotides to the 3' end of the RNA sequence

where the promoter is heterologous with respect to the structural DNA sequence and adapted to cause sufficient expression of the fusion polypeptide to enhance the glyphosate tolerance of a plant cell transformed with said DNA molecule.

In still another aspect of the present invention, a method for selectively controlling weeds in a crop field is presented by planting crop seeds or crop plants transformed with a Class II EPSPS gene to confer glyphosate tolerance to the plants which allows for glyphosate containing herbicides to be applied to the crop to selectively kill the glyphosate sensitive weeds, but not the crops.

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Other and further objects, advantages and aspects of the invention will become apparent from the accompanying drawing figures and the description of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the DNA sequence (SEQ ID NO:1) for the full-length promoter of figwort mosaic virus (FMV35S).

Figure 2 shows the cosmid cloning vector pMON17020.

Figure 3 shows the structural DNA sequence (SEQ ID NO:2) for the Class II EPSPS gene from bacterial isolate Agrobacterium sp. strain CP4 and the deduced amino acid sequence (SEQ ID NO:3).

Figure 4 shows the structural DNA sequence (SEQ ID NO:4) for the Class II EPSPS gene from the bacterial isolate Achromobacter sp. strain LBAA and the deduced amino acid sequence (SEQ ID NO:5).

Figure 5 shows the structural DNA sequence (SEQ ID NO:6) for the Class II EPSPS gene from the bacterial isolate *Pseudomonas sp.* strain PG2982 and the deduced amino acid sequence (SEQ ID NO:7).

Figure 6 shows the Bestfit comparison of the *E. coli* EPSPS amino acid sequence (SEQ ID NO:8) with that for the CP4 EPSPS (SEQ ID NO:3).

Figure 7 shows the Bestfit comparison of the CP4 EPSPS amino acid sequence (SEQ ID NO:3) with that for the LBAA EPSPS (SEQ ID NO:5).

Figure 8 shows the structural DNA sequence (SEQ ID NO:9) for the synthetic CP4 Class II EPSPS gene.

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Figure 9 shows the DNA sequence (SEQ ID NO:10) of the chloroplast transit peptide (CTP) and encoded amino acid sequence (SEQ ID NO:11) derived from the *Arabidopsis thaliana* EPSPS CTP and containing a *SphI* restriction site at the chloroplast processing site, hereinafter referred to as CTP2.

Figure 10 shows the DNA sequence (SEQ ID NO:12) of the chloroplast transit peptide and encoded amino acid sequence (SEQ ID NO:13) derived from the *Arabidopsis thaliana* EPSPS gene and containing an *Eco*RI restriction site within the mature region of the EPSPS, hereinafter referred to as CTP3.

Figure 11 shows the DNA sequence (SEQ ID NO:14) of the chloroplast transit peptide and encoded amino acid sequence (SEQ ID NO:15) derived from the *Petunia hybrida* EPSPS CTP and containing a *SphI* restriction site at the chloroplast processing site and in which the amino acids at the processing site are changed to -Cys-Met-, hereinafter referred to as CTP4.

Figure 12 shows the DNA sequence (SEQ ID NO:16) of the chloroplast transit peptide and encoded amino acid sequence (SEQ ID NO:17) derived from the *Petunia hybrida* EPSPS gene with the naturally occurring *Eco*RI site in the mature region of the EPSPS gene, hereinafter referred to as CTP5.

Figure 13 shows a plasmid map of CP4 plant transformation/expression vector pMON17110.

Figure 14 shows a plasmid map of CP4 synthetic EPSPS gene plant transformation/expression vector pMON17131.

Figure 15 shows a plasmid map of CP4 EPSPS free DNA plant transformation expression vector pMON13640.

Figure 16 shows a plasmid map of CP4 plant transformation/direct selection vector pMON17227.

Figure 17 shows a plasmid map of CP4 plant transformation/expression vector pMON19653.

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STATEMENT OF THE INVENTION

The expression of a plant gene which exists in double-stranded DNA form involves synthesis of messenger RNA (mRNA) from one strand of the DNA by RNA polymerase enzyme, and the subsequent processing of the mRNA primary transcript inside the nucleus. This processing involves a 3' non-translated region which adds polyadenylate nucleotides to the 3' end of the RNA.

Transcription of DNA into mRNA is regulated by a region of DNA usually referred to as the "promoter." The promoter region contains a sequence of bases that signals RNA polymerase to associate with the DNA, and to initiate the transcription into mRNA using one of the DNA strands as a template to make a corresponding complementary strand of RNA.

A number of promoters which are active in plant cells have been described in the literature. These include the nopaline synthase (NOS) and octopine synthase (OCS) promoters (which are carried on tumor-inducing plasmids of Agrobacterium tumefaciens), the cauliflower mosaic virus (CaMV) 19S and 35S promoters, the light-inducible promoter from the small subunit of ribulose bis-phosphate carboxylase (ssRUBISCO, a very abundant plant polypeptide) and the full-length transcript promoter from the figwort mosaic virus (FMV35S). All of these promoters have been used to create various types of DNA constructs which have been expressed in plants; see, e.g., PCT publication WO 84/02913 (Rogers et al., Monsanto).

Promoters which are known or are found to cause transcription of DNA in plant cells can be used in the present invention. Such promoters may be obtained from a variety of sources such as plants and plant DNA viruses and include, but

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are not limited to, the CaMV35S and FMV35S promoters and promoters isolated from plant genes such as ssRUBISCO genes. As described below, it is preferred that the particular promoter selected should be capable of causing sufficient expression to result in the production of an effective amount of a Class II EPSPS to render the plant substantially tolerant to glyphosate herbicides. The amount of Class II EPSPS needed to induce the desired tolerance may vary with the plant species. It is preferred that the promoters utilized have relatively high expression in all meristematic tissues in addition to other tissues inasmuch as it is now known that glyphosate is translocated and accumulated in this type of plant tissue. Alternatively, a combination of chimeric genes can be used to cumulatively result in the necessary overall expression level of the selected Class II EPSPS enzyme to result in the glyphosate tolerant phenotype.

The mRNA produced by a DNA construct of the present invention also contains a 5' non-translated leader sequence. This sequence can be derived from the promoter selected to express the gene, and can be specifically modified so as to increase translation of the mRNA. The 5' non-translated regions can also be obtained from viral RNAs, from suitable eukaryotic genes, or from a synthetic gene sequence. The present invention is not limited to constructs, as presented in the following examples, wherein the non-translated region is derived from both the 5' non-translated sequence that accompanies the promoter sequence and part of the 5' non-translated region of the virus coat protein gene. Rather, the non-translated leader sequence can be derived from an unrelated promoter or coding sequence as discussed above.

A preferred promoter for use in the present invention is the full-length transcript (SEQ ID NO:1) promoter from the

figwort mosaic virus (FMV35S) which functions as a strong and uniform promoter with particularly good expression in meristematic tissue for chimeric genes inserted into plants, particularly dicotyledons. The resulting transgenic plant in general expresses the protein encoded by the inserted gene at a higher and more uniform level throughout the tissues and cells of the transformed plant than the same gene driven by an enhanced CaMV35S promoter. Referring to Figure 1, the DNA sequence (SEQ ID NO:1) of the FMV35S promoter is located between nucleotides 6368 and 6930 of the FMV genome. A 5' non-translated leader sequence is preferably coupled with the promoter. The leader sequence can be from the FMV35S genome itself or can be from a source other than FMV35S.

The 3' non-translated region of the chimeric plant gene contains a polyadenylation signal which functions in plants to cause the addition of polyadenylate nucleotides to the 3' end of the viral RNA. Examples of suitable 3' regions are (1) the 3' transcribed, non-translated regions containing the polyadenylated signal of Agrobacterium tumor-inducing (Ti) plasmid genes, such as the nopaline synthase (NOS) gene, and (2) plant genes like the soybean storage protein genes and the small subunit of the ribulose-1,5-bisphosphate carboxylase (ssRUBISCO) gene. An example of a preferred 3' region is that from the ssRUBISCO gene from pea (E9), described in greater detail below.

The DNA constructs of the present invention also contain a structural coding sequence in double-stranded DNA form which encodes a glyphosate tolerant, highly efficient Class II EPSPS enzyme.

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Identification of glyphosate tolerant, highly efficient EPSPS enzymes

In an attempt to identify and isolate glyphosate tolerant, highly efficient EPSPS enzymes, kinetic analysis of the EPSPS enzymes from a number of bacteria exhibiting tolerance to glyphosate or that had been isolated from suitable sources was undertaken. It was discovered that in some cases the EPSPS enzymes showed no tolerance to inhibition by glyphosate and it was concluded that the tolerance phenotype of the bacterium was due to an impermeability to glyphosate or other factors. In a number of cases, however, microorganisms were identified whose EPSPS enzyme showed a greater degree of tolerance to inhibition by glyphosate and that displayed a low K_m for PEP when compared to that previously reported for other microbial and plant sources. The EPSPS enzymes from these microorganisms were then subjected to further study and analysis.

Table I displays the data obtained for the EPSPS enzymes identified and isolated as a result of the above described analysis. Table I includes data for three identified Class II EPSPS enzymes that were observed to have a high tolerance to inhibition to glyphosate and a low K_m for PEP as well as data for the native Petunia EPSPS and a glyphosate tolerant variant of the Petunia EPSPS referred to as GA101. The GA101 variant is so named because it exhibits the substitution of an alanine residue for a glycine residue at position 101 (with respect to Petunia) in the invariant region. When the change introduced into the Petunia EPSPS (GA101) was introduced into a number of other EPSPS enzymes, similar changes in kinetics were observed, an elevation of the K_i for glyphosate and of the K_m for PEP.

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Table I Kinetic characterization of EPSPS enzymes

5	ENZYME SOURCE	K_m PEP (μM)	K_i Glyphosate (μM)	K _i /K _m
	Petunia	5 200	0.4 2000	0.08 10
	Petunia GA101 PG2982	2.1-3.1 ¹	25-82	~8-4 0
10	LBAA	-7.3-82	60 (est)	~7.9
10	CP4	12 ³	2720	227

- Range of PEP tested = $1-40 \mu M$
- 2 Range of PEP tested = $5-80 \mu M$
- 3 Range of PEP tested = $1.5-40 \mu M$

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The Agrobacterium sp. strain CP4 was initially identified by its ability to grow on glyphosate as a carbon source (10 mM) in the presence of 1 mM phosphate. The strain CP4 was identified from a collection obtained from a fixed-bed immobilized cell column that employed Mannville R-635 diatomaceous earth The column had been run for three months on a waste-water feed from a glyphosate production plant. The column contained 50 mg/ml glyphosate and NH3 as NH4Cl. Total organic carbon was 300 mg/ml and BOD's (Biological Oxygen Demand - a measure of "soft" carbon availability) were less than 30 mg/ml. This treatment column has been described (Heitkamp et al., 1990). Dworkin-Foster minimal salts medium containing glyphosate at 10 mM and with phosphate at 1 mM was used to select for microbes from a wash of this column that were capable of growing on glyphosate as sole carbon source. Dworkin-Foster minimal medium was made up by combining in 1 liter (with autoclaved

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H₂O), 1 ml each of A, B and C and 10 ml of D (as per below) and thiamine HCl (5 mg).

A. D-F Salts (1000X stock; per 100 ml; autoclaved):

5 1 mg H_3BO_3 MnSO₄.7H₂O 1 mg $ZnSO_4.7H_2O$ 12.5 mg CuSO₄.5H₂O 8 mg NaMoO3.3H2O 1.7 mg

> FeSO_{4.7}H₂O (1000X stock; per 100 ml; autoclaved) B.

> > $0.1\,\mathrm{g}$

MgSO₄.7H₂O (1000X stock; per 100 ml; autoclaved) 15 C.

20 g

D. $(NH_4)_2SO_4$ (100X stock; per 100 ml; autoclaved)

20 g

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Yeast Extract (YE; Difco) was added to a final concentration of 0.01 or 0.001%. The strain CP4 was also grown on media composed of D-F salts, amended as described above, containing glucose, gluconate and citrate (each at 0.1 %) as carbon sources and with inorganic phosphate (0.2 - 1.0 mM) as the phosphorous source.

Other Class II EPSPS containing microorganisms were identified as Achromobacter sp. strain LBAA, which was from a collection of bacteria previously described (Hallas et al., 1988), and Pseudomonas sp. strain PG2982 which has been

described in the literature (Moore et al. 1983; Fitzgibbon 1988). It had been reported previously, from measurements in crude lysates, that the EPSPS enzyme from strain PG2982 was less sensitive to inhibition to glyphosate than that of $E.\ coli$, but there has been no report of the details of this lack of sensitivity and there has been no report on the K_m for PEP for this enzyme or of the DNA sequence for the gene for this enzyme (Fitzgibbon, 1988; Fitzgibbon and Braymer, 1990).

10 Relationship of the Class II EPSPS to those previously studied

All EPSPS proteins studied to date have shown a remarkable degree of homology. For example, bacterial and plant EPSPS's are about 54% identical and with similarity as high as 80%. Within bacterial EPSPS's and plant EPSPS's themselves the degree of identity and similarity is much greater (see Table II).

Table II Comparison between exemplary Class I EPSPS
protein sequences!

20	similarity identity				
	E. coli vs. S. typhimurium	93.0	88.3		
	P. hybrida vs. E. coli	71.9	54.5		
	P. hybrida vs. Tomato	92.8	88.2		

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- The EPSPS sequences compared here were obtained from the following references: E. coli, Rogers et al., 1983; S. typhimurium, Stalker et al., 1985; Petunia hybrida, Shah et al., 1986; and Tomato, Gasser et al., 1988.
- When crude extracts of CP4 and LBAA bacteria (50 μg protein) were probed using rabbit anti-EPSPS antibody (Padgette et

al. 1987) to the Petunia EPSPS protein in a Western analysis, no positive signal could be detected, even with extended exposure times (Protein A - ¹²⁵I development system) and under conditions where the control EPSPS (Petunia EPSPS, 20 ng; a Class I EPSPS) was readily detected. The presence of EPSPS activity in these extracts was confirmed by enzyme assay. This surprising result, indicating a lack of similarity between the EPSPS's from these bacterial isolates and those previously studied, coupled with the combination of a low K_m for PEP and a high K_i for glyphosate, illustrates that these new EPSPS enzymes are different from known EPSPS enzymes (now referred to as Class I EPSPS).

Glyphosate Tolerant Enzymes in Microbial Isolates

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For clarity and brevity of disclosure, the following description of the isolation of genes encoding Class II EPSPS enzymes is directed to the isolation of such a gene from a bacterial isolate. Those skilled in the art will recognize that the same or similar strategy can be utilized to isolate such genes from other microbial isolates, plant or fungal sources.

Cloning of the Agrobacterium sp. strain CP4 EPSPS Gene(s) in E. coli

Having established the existence of a suitable EPSPS in Agrobacterium sp. strain CP4, two parallel approaches were undertaken to clone the gene: cloning based on the expected phenotype for a glyphosate tolerant EPSPS; and purification of the enzyme to provide material to raise antibodies and to obtain amino acid sequences from the protein to facilitate the verification of clones. Cloning and genetic techniques, unless otherwise indicated, are generally those described in Maniatis et al., 1982 or

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Sambrook et al., 1987. The cloning strategy was as follows: introduction of a cosmid bank of strain *Agrobacterium* sp. strain CP4 into *E. coli* and selection for the EPSPS gene by selection for growth on inhibitory concentrations of glyphosate.

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Chromosomal DNA was prepared from strain Agrobacterium sp. strain CP4 as follows: The cell pellet from a 200 ml L-Broth (Miller, 1972), late log phase culture of Agrobacterium sp. strain CP4 was resuspended in 10 ml of Solution I; 50 mM Glucose, 10 mM EDTA, 25 mM Tris -CL pH 8.0 (Birnboim and Doly, 1979). SDS was added to a final concentration of 1% and the suspension was subjected to three freeze-thaw cycles, each consisting of immersion in dry ice for 15 minutes and in water at 70°C for 10 minutes. The lysate was then extracted four times with equal volumes of phenol:chloroform (1:1; phenol saturated with TE; TE = 10 mM Tris pH8.0; 1.0 mM EDTA) and the phases separated by centrifugation (15000g; 10 minutes). The ethanol-precipitable material was pelleted from the supernatant by brief centrifugation (8000g; 5 minutes) following addition of two volumes of ethanol. The pellet was resuspended in 5 ml TE and dialyzed for 16 hours at 4°C against 2 liters TE. This preparation yielded a 5 ml DNA solution of 552 μ g/ml.

Partially-restricted DNA was prepared as follows. Three 100 µg aliquot samples of CP4 DNA were treated for 1 hour at 37°C with restriction endonuclease HindIII at rates of 4, 2 and 1 enzyme unit/µg DNA, respectively. The DNA samples were pooled, made 0.25 mM with EDTA and extracted with an equal volume of phenol:chloroform. Following the addition of sodium acetate and ethanol, the DNA was precipitated with two volumes of ethanol and pelleted by centrifugation (12000 g; 10 minutes). The dried DNA pellet was resuspended in 500 µl TE and layered on a 10-40% Sucrose gradient (in 5% increments of 5.5 ml each) in 0.5 M

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NaCl, 50 mM Tris pH8.0, 5 mM EDTA. Following centrifugation for 20 hours at 26,000 rpm in a SW28 rotor, the tubes were punctured and ~1.5 ml fractions collected. Samples (20 μ l) of each second fraction were run on 0.7% agarose gel and the size of the DNA determined by comparison with linearized lambda DNA and HindIII-digested lambda DNA standards. Fractions containing DNA of 25-35 kb fragments were pooled, desalted on AMICON10 columns (7000 rpm; 20°C; 45 minutes) and concentrated by precipitation. This procedure yielded 15 µg of CP4 DNA of the required size. A cosmid bank was constructed using the vector pMON17020. This vector, a map of which is presented in Figure 2, is based on the pBR327 replicon and contains the spectinomycin/streptomycin (Spr;spc) resistance gene from Tn7 (Fling et al., 1985), the chloramphenical resistance gene (Cmr;cat) from Tn9 (Alton et al., 1979), the gene 10 promoter region from phage T7 (Dunn et al., 1983), and the 1.6 kb BglII phage lambda cos fragment from pHC79 (Hohn and Collins, 1980). A number of cloning sites are located downstream of the cat gene. Since the predominant block to the expression of genes from other microbial sources in E. coli appears to be at the level of transcription, the use of the T7 promoter and supplying the T7 polymerase in trans from the pGP1-2 plasmid (Tabor and Richardson, 1985), enables the expression of large DNA segments of foreign DNA, even those containing RNA polymerase transcription termination sequences. The expression of the spc gene is impaired by transcription from the T7 promoter such that only Cmr can be selected in strains containing pGP1-2. The use of antibiotic resistances such as Cm resistance which do not employ a membrane component is preferred due to the observation that high level expression of resistance genes that involve a membrane component, i.e. B-lactamase and Amp resistance, give rise to a glyphosate tolerant

phenotype. Presumably, this is due to the exclusion of glyphosate from the cell by the membrane localized resistance protein. It is also preferred that the selectable marker be oriented in the same direction as the T7 promoter.

The vector was then cut with *HindIII* and treated with calf alkaline phosphatase (CAP) in preparation for cloning. Vector and target sequences were ligated by combining the following:

10	Vector DNA (HindIII/CAP)	3 µg
	Size fractionated CP4 HindIII fragments	1.5 µg
	10X ligation buffer	2.2 µl
	T4 DNA ligase (New England Biolabs) (400 U/µl)	1.0 யி

15 and adding H₂O to 22.0 μl. This mixture was incubated for 18 hours at 16°C. 10X ligation buffer is 250 mM Tris-HCl, pH 8.0; 100 mM MgCl₂; 100 mM Dithiothreitol; 2 mM Spermidine. The ligated DNA (5 μl) was packaged into lambda phage particles (Stratagene; Gigapack Gold) using the manufacturer's procedure.

A sample (200 μl) of E. coli HB101 (Boyer and Rolland-Dussoix, 1973) containing the T7 polymerase expression plasmid pGP1-2 (Tabor and Richardson, 1985) and grown overnight in L-Broth (with maltose at 0.2% and kanamycin at 50 μg/ml) was infected with 50 μl of the packaged DNA. Transformants were selected at 30°C on M9 (Miller, 1972) agar containing kanamycin (50 μg/ml), chloramphenicol (25 μg/ml), L-proline (50 μg/ml), L-leucine (50 μg/ml) and B1 (5 μg/ml), and with glyphosate at 3.0 mM. Aliquot samples were also plated on the same media lacking glyphosate to titer the packaged cosmids.

30 Cosmid transformants were isolated on this latter medium at a

rate of ~5 x 105 per μg CP4 HindIII DNA after 3 days at 30°C. Colonies arose on the glyphosate agar from day 3 until day 15 with a final rate of ~1 per 200 cosmids. DNA was prepared from 14 glyphosate tolerant clones and, following verification of this 5 phenotype, was transformed into E. coli GB100/pGP1-2 (E. coli GB100 is an aroA derivative of MM294 [Talmadge and Gilbert, 1980]) and tested for complementation for growth in the absence of added aromatic amino acids and aminobenzoic acids. Other aroA strains such as SR481 (Bachman et al. 1980; Padgette et al., 1987), 10 could be used and would be suitable for this experiment. The use of GB100 is merely exemplary and should not be viewed in a limiting sense. This aroA strain usually requires that growth media be supplemented with L-phenylalanine, L-tyrosine and L-tryptophan each at 100 µg/ml and with para-hydroxybenzoic 15 acid, 2,3-dihydroxybenzoic acid and para-aminobenzoic acid each at 5 µg/ml for growth in minimal media. Of the fourteen cosmids tested only one showed complementation of the aroA- phenotype. Transformants of this cosmid, pMON17076, showed weak but uniform growth on the unsupplemented minimal media after 10 20 days.

The proteins encoded by the cosmids were determined in vivo using a T7 expression system (Tabor and Richardson, 1985). Cultures of E. coli containing pGP1-2 (Tabor and Richardson, 1985) and test and control cosmids were grown at 30°C in L-broth (2 ml) with chloramphenicol and kanamycin (25 and 50 μg/ml, respectively) to a Klett reading of ~ 50. An aliquot was removed and the cells collected by centrifugation, washed with M9 salts (Miller, 1972) and resuspended in 1 ml M9 medium containing glucose at 0.2%, thiamine at 20 μg/ml and containing 30 the 18 amino acids at 0.01% (minus cysteine and methionine). Following incubation at 30°C for 90 minutes, the cultures were

transferred to a 42°C water bath and held there for 15 minutes. Rifampicin (Sigma) was added to 200 μg/ml and the cultures held at 42°C for 10 additional minutes and then transferred to 30°C for 20 minutes. Samples were pulsed with 10 μCi of 35S-methionine for 5 minutes at 30°C. The cells were collected by centrifugation and suspended in 60-120 μl cracking buffer (60 mM Tris-HCl 6.8, 1% SDS, 1% 2-mercaptoethanol, 10% glycerol, 0.01% bromophenol blue). Aliquot samples were electrophoresed on 12.5% SDS-PAGE and following soaking for 60 minutes in 10 volumes of Acetic 10 Acid-Methanol-water (10:30:60), the gel was soaked in ENLIGHTNING TM (DUPONT) following manufacturer's directions, dried, and exposed at -70°C to X-Ray film. Proteins of about 45 kd in size, labeled with 35S-methionine, were detected in number of the cosmids, including pMON17076.

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Purification of EPSPS from Agrobacterium sp. strain CP4

All protein purification procedures were carried out at 3-5°C. EPSPS enzyme assays were performed using either the phosphate release or radioactive HPLC method, as previously described in Padgette et al. 1987, using 1 mM phosphoenol pyruvate (PEP, Boehringer) and 2 mM shikimate-3-phosphate (S3P) substrate concentrations. For radioactive HPLC assays, 14C-PEP (Amersham) was utilized. S3P was synthesized as previously described in Wibbenmeyer et al. 1988. N-terminal amino acid sequencing was performed by loading samples onto a Polybrene precycled filter in aliquots while drying. Automated Edman degradation chemistry was used to determine the N-terminal protein sequence, using an Applied Biosytems Model 470A gas phase sequencer (Hunkapiller et al. 1983) with an 30 Applied Biosystems 120A PTH analyzer.

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Five 10-litre fermentations were carried out on a spontaneous "smooth" isolate of strain CP4 that displayed less clumping when grown in liquid culture. This reduced clumping and smooth colony morphology may be due to reduced polysaccharide production by this isolate. In the following section dealing with the purification of the EPSPS enzyme, CP4 refers to the "smooth" isolate - CP4-S1. The cells from the three batches showing the highest specific activities were pooled. Cell paste of Agrobacterium sp. CP4 (300 g) was washed twice with 0.5 L of 0.9% 10 saline and collected by centrifugation (30 minutes, 8000 rpm in a GS3 Sorvall rotor). The cell pellet was suspended in 0.9 L extraction buffer (100 mM TrisCl, 1 mM EDTA, 1 mM BAM (Benzamidine), 5 mM DTT, 10% glycerol, pH 7.5) and lysed by 2 passes through a Manton Gaulin cell. The resulting solution was 15 centrifuged (30 minutes, 8000 rpm) and the supernatant was treated with 0.21 L of 1.5% protamine sulfate (in 100 mM TrisCl, pH 7.5, 0.2% w/v final protamine sulfate concentration). After stirring for 1 hour, the mixture was centrifuged (50 minutes, 8000 rpm) and the resulting supernatant treated with solid ammonium 20 sulfate to 40% saturation and stirred for 1 hour. centrifugation (50 minutes, 8000 rpm), the resulting supernatant was treated with solid ammonium sulfate to 70% saturation, stirred for 50 minutes, and the insoluble protein was collected by centrifugation (1 hour, 8000 rpm). This 40-70% ammonium sulfate 25 fraction was then dissolved in extraction buffer to give a final volume of 0.2 L, and dialyzed twice (Spectrum 10,000 MW cutoff dialysis tubing) against 2 L of extraction buffer for a total of 12 hours.

To the resulting dialyzed 40-70% ammonium sulfate 30 fraction (0.29 L) was added solid ammonium sulfate to give a final concentration of 1 M. This material was loaded (2 ml/min) onto a column (5 cm x 15 cm, 295 ml) packed with phenyl Sepharose CL-4B (Pharmacia) resin equilibrated with extraction buffer containing 1 M ammonium sulfate, and washed with the same buffer (1.5 L, 2 ml/min). EPSPS was eluted with a linear gradient of extraction buffer going from 1 M to 0.00 M ammonium sulfate (total volume of 1.5 L, 2 ml/min). Fractions were collected (20 ml) and assayed for EPSPS activity by the phosphate release assay. The fractions with the highest EPSPS activity (fractions 36-50) were pooled and dialyzed against 3 x 2 L (18 hours) of 10 mM TrisCl, 25 mM KCl, 1 mM EDTA, 5 mM DTT, 10% glycerol, pH 7.8.

The dialyzed EPSPS extract (350 ml) was loaded (5 ml/min) onto a column (2.4 cm x 30 cm, 136 ml) packed with Q-Sepharose Fast Flow (Pharmacia) resin equilibrated with 10 mM TrisCl, 25 mM KCl, 5 mM DTT, 10% glycerol, pH 7.8 (Q Sepharose buffer), and washed with 1 L of the same buffer. EPSPS was eluted with a linear gradient of Q Sepharose buffer going from 0.025 M to 0.40 M KCl (total volume of 1.4 L, 5 ml/min). Fractions were collected (15 ml) and assayed for EPSPS activity by the phosphate release assay. The fractions with the highest EPSPS activity (fractions 47-60) were pooled and the protein was precipitated by adding solid ammonium sulfate to 80% saturation and stirring for 1 hour. The precipitated protein was collected by centrifugation (20 minutes, 12000 rpm in a GSA Sorvall rotor), dissolved in Q Sepharose buffer (total volume of 14 ml), and dialyzed against the same buffer (2 x 1 L, 18 hours).

The resulting dialyzed partially purified EPSPS extract (19 ml) was loaded (1.7 ml/min) onto a Mono Q 10/10 column (Pharmacia) equilibrated with Q Sepharose buffer, and washed with the same buffer (35 ml). EPSPS was eluted with a linear gradient of 0.025 M to 0.35 M KCl (total volume of 119 ml, 1.7 ml/min). Fractions were collected (1.7 ml) and assayed for EPSPS

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activity by the phosphate release assay. The fractions with the highest EPSPS activity (fractions 30-37) were pooled (6 ml).

The Mono Q pool was made 1 M in ammonium sulfate by the addition of solid ammonium sulfate and 2 ml aliquots were chromatographed on a Phenyl Superose 5/5 column (Pharmacia) equilibrated with 100 mM TrisCl, 5 mM DTT, 1 M ammonium sulfate, 10% glycerol, pH 7.5 (Phenyl Superose buffer). Samples were loaded (1 ml/min), washed with Phenyl Superose buffer (10 ml), and eluted with a linear gradient of Phenyl Superose buffer going from 1 M to 0.00 M ammonium sulfate (total volume of 60 ml, 1 ml/min). Fractions were collected (1 ml) and assayed for EPSPS activity by the phosphate release assay. The fractions from each run with the highest EPSPS activity (fractions ~36-40) were pooled together (10 ml, 2.5 mg protein). For N-terminal amino acid sequence determination, a portion of one fraction (#39 from run 1) was dialyzed against 50 mM NaHCO₃ (2 x 1 L). resulting pure EPSPS sample (0.9 ml, 77 µg protein) was found to exhibit a single N-terminal amino acid sequence of: XH(G)ASSRPATARKSS(G)LX(G)(T)V(R)IPG(D)(K)(M) (SEQ ID NO:18).

In this and all amino acid sequences to follow, the standard single letter nomenclature is used. All peptide structures represented in the following description are shown in conventional format wherein the amino group at the N-terminus appears to the left and the carboxyl group at the C-terminus at the right. Likewise, amino acid nomenclature for the naturally occurring amino acids found in protein is as follows: alanine (Ala;A), asparagine (Asn;N), aspartic acid (Asp;D), arginine (Arg;R), cysteine (Cys;C), glutamic acid (Glu;E), glutamine (Gln;Q), glycine (Gly;G), histidine (His;H), isoleucine (Ile;I), leucine (Leu;L), lysine (Lys;K), methionine (Met;M), phenylalanine (Phe;F), proline (Pro;P), serine (Ser;S), threonine

(Thr;T), tryptophan (Trp;W), tyrosine (Tyr;Y), and valine (Val;V). An "X" is used when the amino acid residue is unknown and parentheses designate that an unambiguous assignment is not possible and the amino acid designation within the parentheses is the most probable estimate based on known information.

The remaining Phenyl Superose EPSPS pool was dialyzed against 50 mM TrisCl, 2 mM DTT, 10 mM KCl, 10% glycerol, pH 7.5 (2 x 1 L). An aliquot (0.55 ml, 0.61 mg protein) was loaded (1 ml/min) onto a Mono Q 5/5 column (Pharmacia) equilibrated with Q Sepharose buffer, washed with the same buffer (5 ml), and eluted with a linear gradient of Q Sepharose buffer going from 0-0.14 M KCl in 10 minutes, then holding at 0.14 M KCl (1 ml/min). Fractions were collected (1 ml) and assayed for EPSPS activity by the phosphate release assay and were subjected to SDSPAGE (10-15%, Phast System, Pharmacia, with silver staining) to determine protein purity. Fractions exhibiting a single band of protein by SDS-PAGE (22-25, 222 μg) were pooled and dialyzed against 100 mM ammonium bicarbonate, pH 8.1 (2 x 1 L, 9 hours).

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Tryspinolysis and peptide sequencing of Agrobacterium sp strain CP4 EPSPS

To the resulting pure Agrobacterium sp. strain CP4 EPSPS (111 μg) was added 3 μg of trypsin (Calbiochem), and the trypsinolysis reaction was allowed to proceed for 16 hours at 37°C. The tryptic digest was then chromatographed (1ml/min) on a C18 reverse phase HPLC column (Vydac) as previously described in Padgette et al. 1988 for E. coli EPSPS. For all peptide purifications, 0.1% trifluoroacetic acid (TFA, Pierce) was designated buffer "RP-A" and 0.1% TFA in acetonitrile was buffer "RP-B". The gradient used for elution of the trypsinized Agrobacterium sp. CP4

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EPSPS was: 0-8 minutes, 0% RP-B; 8-28 minutes, 0-15% RP-B; 28-40 minutes, 15-21% RP-B; 40-68 minutes, 21-49% RP-B; 68-72 minutes, 49-75% RP-B; 72-74 minutes, 75-100% RP-B. Fractions were collected (1 ml) and, based on the elution profile at 210 nm, at least 70 distinct peptides were produced from the trypsinized EPSPS. Fractions 40-70 were evaporated to dryness and redissolved in 150 μl each of 10% acetonitrile, 0.1% trifluoroacetic acid.

The fraction 61 peptide was further purified on the C18 column by the gradient: 0-5 minutes, 0% RP-B; 5-10 minutes, 0-38% RP-B; 10-30 minutes, 38-45% B. Fractions were collected based on the UV signal at 210 nm. A large peptide peak in fraction 24 eluted at 42% RP-B and was dried down, resuspended as described above, and rechromatographed on the C18 column with the gradient: 0-5 minutes, 0% RP-B; 5-12 min, 0-38% RP-B; 12-15 min, 38-39% RP-B; 15-18 minutes, 39% RP-B; 18-20 minutes, 39-41% RP-B; 20-24 minutes, 41% RP-B; 24-28 minutes, 42% RP-B. The peptide in fraction 25, eluting at 41% RP-B and designated peptide 61-24-25, was subjected to N-terminal amino acid sequencing, and the following sequence was determined:

APSM(I)(D)EYPILAV (SEQ ID NO:19).

The CP4 EPSPS fraction 53 tryptic peptide was further purified by C18 HPLC by the gradient 0% B (5 minutes), 0-30% B (5-17 minutes), 30-40% B (17-37 minutes). The peptide in fraction 28, eluting at 34% B and designated peptide 53-28, was subjected to N-terminal amino acid sequencing, and the following sequence was determined:

ITGLLEGEDVINTGK (SEQ ID NO: 20).

In order to verify the CP4 EPSPS cosmid clone, a number of oligonucleotide probes were designed on the basis of the sequence of two of the tryptic sequences from the CP4 enzyme (Table III). The probe identified as MID was very low degeneracy and was used for initial screening. The probes identified as EDV-C and EDV-T were based on the same amino acid sequences and differ in one position (underlined in Table III below) and were used as confirmatory probes, with a positive to be expected only from one of these two probes. In the oligonucleotides below, alternate acceptable nucleotides at a particular position are designated by a "/" such as A/C/T.

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Table III Selected CP4 EPSPS peptide sequences and DNA probes

PEPTIDE 61-24-25 APSM(I)(D)EYPILAV (SEQ ID NO:19)
Probe MID; 17-mer; mixed probe; 24-fold degenerate

ATGATA/C/TGAC/TGAG/ATAC/TCC (SEQ ID NO:21)
PEPTIDE 53-28 ITGLLEGEDVINTGK (SEQ ID NO:20)
Probe EDV-C; 17-mer; mixed probe; 48-fold degenerate
GAA/GGAC/TGTA/C/G/TATA/C/TAACAC (SEQ ID NO:22)
Probe EDV-T; 17-mer; mixed probe; 48-fold degenerate
GAA/GGAC/TGTA/C/G/TATA/C/TAATAC (SEQ ID NO:23)

The probes were labeled using gamma-32P-ATP and polynucleotide kinase. DNA from fourteen of the cosmids described above was restricted with EcoRI, transferred to membrane and probed with the olignucleotide probes. The conditions used were as follows: prehybridization was carried out in 6X SSC, 10X Denhardt's for 2-18 hour periods at 60°C, and hybridization was for 48-72 hours in 6X SSC, 10X Denhardt's, 100 µg/ml tRNA at 10°C below the T_d for the probe. The T_d of the probe was approximated by the formula 2°C x (A+T) + 4°C x (G+C). The

filters were then washed three times with 6X SSC for ten minutes each at room temperature, dried and autoradiographed. Using the MID probe, an ~9.9 kb fragment in the pMON17076 cosmid gave the only positive signal. This cosmid DNA was then probed with the EDV-C (SEQ ID NO:22) and EDV-T (SEQ ID NO:23) probes separately and again this ~9.9 kb band gave a signal and only with the EDV-T probe.

The combined data on the glyphosate tolerant phenotype, the complementation of the E. coli aroA- phenotype, the expression of a ~45 Kd protein, and the hybridization to two probes derived from the CP4 EPSPS amino acid sequence strongly suggested that the pMON17076 cosmid contained the EPSPS gene.

Localization and subcloning of the CP4 EPSPS gene

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The CP4 EPSPS gene was further localized as follows: a number of additional Southern analyses were carried out on different restriction digests of pMON17076 using the MID (SEQ ID NO:21) and EDV-T (SEQ ID NO:23) probes separately. Based on these analyses and on subsequent detailed restriction mapping of 20 the pBlueScript (Stratagene) subclones of the ~9.9 kb fragment from pMON17076, a 3.8 kb EcoRI-SalI fragment was identified to which both probes hybridized. This analysis also showed that MID (SEQ ID NO:21) and EDV-T (SEQ ID NO:23) probes hybridized to different sides of BamHI, ClaI, and SacII sites. This 3.8 kb 25 fragment was cloned in both orientations in pBlueScript to form pMON17081 and pMON17082. The phenotypes imparted to E. coli by these clones were then determined. Glyphosate tolerance was determined following transformation into E. coli MM294 containing pGP1-2 (pBlueScript also contains a T7 promoter) on M9 agar media containing glyphosate at 3 mM. Both pMON17081 and pMON17082 showed glyphosate tolerant colonies at three days

at 30°C at about half the size of the controls on the same media lacking glyphosate. This result suggested that the 3.8 kb fragment contained an intact EPSPS gene. The apparent lack of orientation-dependence of this phenotype could be explained by the presence of the T7 promoter at one side of the cloning sites and the lac promoter at the other. The aroA phenotype was determined in transformants of E. coli GB100 on M9 agar media lacking aromatic supplements. In this experiment, carried out with and without the Plac inducer IPTG, pMON17082 showed much greater growth than pMON17081, suggesting that the EPSPS gene was expressed from the SalI site towards the EcoRI site.

Nucleotide sequencing was begun from a number of restriction site ends, including the BamHI site discussed above. Sequences encoding protein sequences that closely matched the N-terminus protein sequence and that for the tryptic fragment 53-28 (SEQ ID NO:20) (the basis of the EDV-T probe) (SEQ ID NO:23)were localized to the SalI side of this BamHI site. These data provided conclusive evidence for the cloning of the CP4 EPSPS gene and for the direction of transcription of this gene. These data coupled with the restriction mapping data also 20 indicated that the complete gene was located on an ~2.3 kb XhoI fragment and this fragment was subcloned into pBlueScript. The sequence of almost 2 kb of this fragment was nucleotide combination of sequencing from cloned determined by a restriction fragments and by the use of specific primers to extend the sequence. The nucleotide sequence of the CP4 EPSPS gene and flanking regions is shown in Figure 3 (SEQ ID NO:2). The sequence corresponding to peptide 61-24-25 (SEQ ID NO:19) was also located. The sequence was determined using both the Sequenase kit from IBI (International Biotechnologies Inc.) and the T7 sequencing /Deaza Kit from Pharmacia.

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That the cloned gene encoded the EPSPS activity purified from the Agrobacterium sp. strain CP4 was verified in the following manner: By a series of site directed mutageneses, $Bgl\Pi$ and NcoI sites were placed at the N-terminus with the fMet contained within the NcoI recognition sequence, the first internal NcoI site was removed (the second internal NcoI site was removed later), and a SacI site was placed after the stop codons. At a later stage the internal NotI site was also removed by site-directed mutagenesis. The following list includes the primers for the 10 site-directed mutagenesis (addition or removal of restriction sites) of the CP4 EPSPS gene. Mutagenesis was carried out by the procedures of Kunkel et al. (1987), essentially as described in Sambrook et al. (1989).

15 PRIMER BgNc (addition of BglII and Ncol sites to N-terminus) CGTGGATAGATCTAGGAAGACAACCATGGCTCACGGTC (SEQ ID NO:24)

PRIMER Sph2 (addition of SphI site to N-terminus) 20 GGATAGATTAAGGAAGACGCGCATGCTTCACGGTGCAAGC AGCC (SEQ ID NO:25)

PRIMER S1 (addition of SacI site immediately after stop codons) GGCTGCCTGATGAGCTCCACAATCGCCATCGATGG 25 (SEQ ID NO:26)

PRIMER N1 (removal of internal NotI recognition site) CGTCGCTCGTCGTGCGTGGCCCCTGACGGC (SEQ ID NO:27)

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PRIMER Nco1 (removal of first internal NcoI recognition site)
CGGGCAAGGCCATGCAGGCTATGGGCGCC (SEQ ID NO:28)

PRIMER Nco2 (removal of second internal NcoI recognition site)
CGGGCTGCCGCCTGACTATGGGCCTCGTCGG (SEQ ID NO:29)

This CP4 EPSPS gene was then cloned as a NcoI-BamHI N-terminal fragment plus a BamHI-SacI C-terminal fragment into a PrecA-gene 10L expression vector similar to those described (Wong et al., 1988; Olins et al., 1988) to form pMON17101. The K_m for PEP and the K_i for glyphosate were determined for the EPSPS activity in crude lysates of pMON17101/GB100 transformants following induction with nalidixic acid (Wong et al., 1988) and found to be the same as that determined for the purified and crude enzyme preparations from Agrobacterium sp. strain CP4.

Characterization of the EPSPS gene from Achromobacter sp. strain LBAA and from Pseudomonas sp. strain PG2982

A cosmid bank of partially *HindIII*-restricted LBAA DNA was constructed in *E. coli* MM294 in the vector pHC79 (Hohn and Collins, 1980). This bank was probed with a full length CP4 EPSPS gene probe by colony hybridization and positive clones were identified at a rate of ~1 per 400 cosmids. The LBAA EPSPS gene was further localized in these cosmids by Southern analysis. The gene was located on an ~2.8 kb *XhoI* fragment and by a series of sequencing steps, both from restriction fragment ends and by using the oligonucleotide primers from the sequencing of the CP4 EPSPS gene, the nucleotide sequence of the LBAA EPSPS gene was completed and is presented in Figure 4 (SEQ ID NO:4).

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The EPSPS gene from PG2982 was also cloned. The EPSPS protein was purified, essentially as described for the CP4 enzyme, with the following differences: Following the Sepharose CL-4B column, the fractions with the highest EPSPS activity were pooled and the protein precipitated by adding solid ammonium sulfate to 85% saturation and stirring for 1 hour. The precipitated protein was collected by centrifugation, resuspended in Q Sepharose buffer and following dialysis against the same buffer was loaded onto the column (as for the CP4 enzyme). After purification on the Q Sepharose column, ~40 mg of protein in 100 mM Tris pH 7.8, 10% glycerol, 1 mM EDTA, 1 mM DTT, and 1 M ammonium sulfate, was loaded onto a Phenyl Superose (Pharmacia) column. The column was eluted at 1.0 ml/minutes with a 40 ml gradient from 1.0 M to 0.00 M ammonium sulfate in the above buffer.

Approximately 1.0 mg of protein from the active fractions of the Phenyl Superose 10/10 column was loaded onto a Pharmacia Mono P 5/10 Chromatofocusing column with a flow rate of 0.75 ml/minutes. The starting buffer was 25 mM bis-Tris at pH 6.3, and the column was eluted with 39 ml of Polybuffer 74, pH 4.0. Approximately 50 µg of the peak fraction from the Chromatofocusing column was dialyzed into 25 mM ammonium bicarbonate. This sample was then used to determine the N-terminal amino acid sequence.

The N-terminal sequence obtained was:

XHSASPKPATARRSE (where X = an unidentified residue) (SEQ ID NO:30). A number of degenerate oligonucleotide probes were designed based on this sequence and used to probe a library of PG2982 partial-HindIII DNA in the cosmid pHC79 (Hohn and Collins, 1980) by colony hybridization under nonstringent conditions. Final washing conditions were 15

minutes with 1X SSC, 0.1% SDS at 55°C. One probe with the sequence GCGGTBGCSGGYTTSGG (where B = C, G, or T; S = C or G, and Y = C or T) (SEQ ID NO:31) identified a set of cosmid clones.

The cosmid set identified in this way was made up of cosmids of diverse *Hind*III fragments. However, when this set was probed with the CP4 EPSPS gene probe, a cosmid containing the PG2982 EPSPS gene was identified (designated as cosmid 9C1 originally and later as pMON20107). By a series of restriction mappings and Southern analysis this gene was localized to a ~2.8 kb *Xho*I fragment and the nucleotide sequence of this gene was determined. This DNA sequence (SEQ ID NO:6) is shown in Figure 5. There are no nucleotide differences between the EPSPS gene sequences from LBAA (SEQ ID NO:4) and PG2982 (SEQ ID NO:6). The kinetic parameters of the two enzymes are within the range of experimental error.

A gene from PG2982 that imparts glyphosate tolerance in *E. coli* has been sequenced (Fitzgibbon, 1988; Fitzgibbon and Braymer, 1990). The sequence of the PG2982 EPSPS Class II gene shows no homology to the previously reported sequence suggesting that the glyphosate tolerant phenotype of the previous work is not related to EPSPS.

Alternative Isolation Protocols for Other Class II EPSPS

25 Structural Genes

A number of Class II genes have been isolated and described here. It is clear that the initial gene cloning, that of the gene from CP4, was difficult due to the low degree of similarity between the Class I and Class II enzymes and genes. The identification of the other genes however was greatly facilitated by the use of this first gene as a probe. In the cloning of the LBAA

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EPSPS gene, the CP4 gene probe allowed the rapid identification of cosmid clones and the localization of the intact gene to a small restriction fragment and some of the CP4 sequencing primers were also used to sequence the LBAA (and PG2982) EPSPS gene(s). The CP4 gene probe was also used to confirm the PG2982 gene clone. The high degree of similarity of the Class II EPSPS genes may be used to identify and clone additional genes in much the same way that Class I EPSPS gene probes have been used to clone other Class I genes. An example of the latter was in the cloning of the A. thaliana EPSPS gene using the P. hybrida gene as a probe (Klee et al., 1987).

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Glyphosate tolerant EPSPS activity has been reported previously for EPSP synthases from a number of sources. These enzymes have not been characterized to any extent in most cases. The use of Class I and Class II EPSPS gene probes or antibody probes provide a rapid means of initially screening for the nature of the EPSPS and provide tools for the rapid cloning and characterization of the genes for such enzymes.

Dacteria that were isolated from a glyphosate treatment facility (Strains CP4 and LBAA). The third (PG2982) was from a bacterium that had been isolated from a culture collection strain. This latter isolation suggests that exposure to glyphosate may not be a prerequisite for the isolation of high glyphosate tolerant EPSPS enzymes and that the screening of collections of bacteria could yield additional isolates. It is possible to enrich for glyphosate degrading or glyphosate resistant microbial populations (Quinn et al., 1988; Talbot et al., 1984) in cases where it was felt that enrichment for such microorganisms would enhance the isolation frequency of Class II EPSPS microorganisms. Additional bacteria containing class II EPSPS gene have also been

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identified. A bacterium called C12, isolated from the same treatment column beads as CP4 (see above) but in a medium in which glyphosate was supplied as both the carbon and phosphorus source, was shown by Southern analysis to hybridize with a probe consisting of the CP4 EPSPS coding sequence. This result, in conjunction with that for strain LBAA, suggests that this enrichment method facilitates the identification of Class II EPSPS isolates. New bacterial isolates containing Class II EPSPS genes also been identified from environments other than glyphosate waste treatment facilities. An inoculum was prepared by extracting soil (from a recently harvested soybean field in Jerseyville, Illinois) and a population of bacteria selected by growth at 28°C in Dworkin-Foster medium containing glyphosate at 10 mM as a source of carbon (and with cycloheximide at 100 µg/ml to prevent the growth of fungi). Upon plating on L-agar media, five colony types were identified. Chromosomal DNA was prepared from 2ml L-broth cultures of these isolates and the presence of a Class II EPSPS gene was probed using a the CP4 EPSPS coding sequence probe by Southern analysis under stringent hybridization and washing conditions. One of the soil isolates, S2, was positive by this screen.

Relationships between different EPSPS genes

The deduced amino acid sequences of a number of Class I and the Class II EPSPS enzymes were compared using the Bestfit computer program provided in the UWGCG package (Devereux et al. 1984). The degree of similarity and identity as determined using this program is reported. The degree of similarity/identity determined within Class I and Class II protein sequences is remarkably high, for instance, comparing *E. coli* with *S. typhimurium* (similarity/identity = 93%/88%) and even

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comparing E. coli with a plant EPSPS (Petunia hybrida; 72%/55%). This data is shown in Table IV. The comparison of sequences between Class I and Class II, however, shows only a very low degree of relatedness between the Classes (similarity/identity = 50-53%/23-30%). The display of the Bestfit analysis for the E.coli (SEQ ID NO:8) and CP4 (SEQ ID NO:3) sequences shows the positions of the conserved residues and is presented in Figure 6. Previous analyses of EPSPS sequences had noted the high degree of conservation of sequences of the enzymes and the almost 10 invariance of sequences in two regions - the "20-35" and "95-107" regions (Gasser et al., 1988; numbered according to the Petunia EPSPS sequence) - and these regions are less conserved in the case of CP4 and LBAA when compared to Class I bacterial and plant EPSPS sequences (see Figure 6 for a comparison of the E. coli and 15 CP4 EPSPS sequences with the E. coli sequence appearing as the top sequence in the Figure). The corresponding sequences in the CP4 Class II EPSPS are: PGDKSISHRSFMFGGL (SEQ ID NO:32) and LDFGNAATGCRLT

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(SEQ ID NO:33).

These comparisons show that the overall relatedness of Class I and Class II is EPSPS proteins is low and that sequences in putative conserved regions have also diverged considerably.

In the CP4 EPSPS an alanine residue is present at the 25 "glycine101" position. The replacement of the conserved glycine (from the "95-107" region) by an alanine results in an elevated K_i for glyphosate and in an elevation in the K_m for PEP in Class I EPSPS. In the case of the CP4 EPSPS, which contains an alanine at this position, the K_m for PEP is in the low range, indicating that the Class II enzymes differ in many aspects from the EPSPS enzymes heretofore characterized.

Within the Class II isolates, the degree of similarity/identity is as high as that noted for that within Class I (Table IV). Figure 7 displays the Bestfit computer program alignment of the CP4 (SEQ ID NO:3) and LBAA (SEQ ID NO:5) EPSPS deduced amino acid sequences with the CP4 sequence appearing as the top sequence in the Figure. The symbols used in Figures 6 and 7 are the standard symbols used in the Bestfit computer program to designate degrees of similarity and identity.

10 Table IV Comparison of relatedness of EPSPS protein sequences Comparison between Class I and Class II EPSPS protein

		sequences	
		<u>similarity</u>	identity
	E. coli vs. CP4	52.8	26.3
15	E. coli vs. LBAA	52.1	26.7
	S. typhimurium vs. CP4	51.8	25.8
	B. pertussis vs. CP4	52.8	27.3
	S. cerevisiae vs. CP4	53.5	29.9
	P. hybrida vs. CP4	50.2	23.4

Comparison between Class I EPSPS protein sequences

	<u>similarity</u>	identity
E. coli vs. S. typhimurium	93.0	88.3
P. hybrida vs. E. coli	71.9	54.5

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Comparison between Class II EPSPS protein sequences

	<u>similarity</u>	identity
Agrobacterium sp. strain CP	4	
vs. Achromobacter sp.		•
strain LBAA	89.9	83.7

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The EPSPS sequences compared here were obtained from the following references: E. coli, Rogers et al., 1983; S. typhimurium, Stalker et al., 1985; Petunia hybrida, Shah et al., 1986; B. pertussis, Maskell et al., 1988; and S. cerevisiae, Duncan et al., 1987.

One difference that may be noted between the deduced amino acid sequences of the CP4 and LBAA EPSPS proteins is at position 100 where an Alanine is found in the case of the CP4 enzyme and a Glycine is found in the case of the LBAA enzyme. In the Class I EPSPS enzymes a Glycine is usually found in the equivalent position, i.e Glycine96 in E. coli and K. pneumoniae and Glycine 101 in Petunia. In the case of these three enzymes it has been reported that converting that Glycine to an Alanine results in an elevation of the appKi for glyphosate and a concomitant elevation in the appKm for PEP (Kishore et al. 1986; Kishore and Shah, 1988; Sost and Amrhein, 1990), which, as discussed above. makes the enzyme less efficient especially under conditions of lower PEP concentrations. The Glycine 100 of the LBAA EPSPS was converted to an Alanine and both the appKm for PEP and the appKi for glyphosate were determined for the variant. Glycine 100 Alanine change was introduced by mutagenesis using the following primer:

and both the wild type and variant genes were expressed in *E. coli* in a *RecA* promoter expression vector (pMON17201 and pMON17264, respectively) and the appKm's and appKi's determined in crude lysates. The data indicate that the appKi(glyphosate) for the G100A variant is elevated about 16-fold (Table V). This result is in agreement with the observation of the importance of this G-A change in raising the appKi(glyphosate) in

the Class I EPSPS enzymes. However, in contrast to the results in the Class I G-A variants, the appKm(PEP) in the Class II (LBAA) G-A variant is unaltered. This provides yet another distinction between the Class II and Class I EPSPS enzymes.

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Table V

		appKm(PEP)	appKi(glyphosate)
	Lysate prepared from:		
10	E. coli/pMON17201 (wild type)	5.3 μM	28 μM*
	E. coli/pMON17264	5.5 μM	459 μM#
	(G100A variant)		

@ range of PEP: 2-40 µM

* range of glyphosate: 0-310 μ M; # range of glyphosate: 0-5000 μ M.

The LBAA G100A variant, by virtue of its superior kinetic properties, is capable of imparting improved glyphosate in planta.

20 <u>Modification and Resynthesis of the Agrobacterium sp. strain CP4</u> EPSPS Gene Sequence

The EPSPS gene from Agrobacterium sp. strain CP4 contains sequences that could be inimical to high expression of the gene in plants. These sequences include potential polyadenylation sites that are often and A+T rich, a higher G+C% than that frequently found in plant genes (63% versus ~50%), concentrated stretches of G and C residues, and codons that are not used frequently in plant genes. The high G+C% in the CP4 EPSPS gene has a number of potential consequences including the following: a higher usage of G or C than that found in plant genes in the third position in codons, and the potential to form strong hair-pin

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structures that may affect expression or stability of the RNA. The reduction in the G+C content of the CP4 EPSPS gene, the disruption of stretches of G's and C's, the elimination of potential polyadenylation sequences, and improvements in the codon usage to that used more frequently in plant genes, could result in higher expression of the CP4 EPSPS gene in plants.

A synthetic CP4 gene was designed to change as completely as possible those inimical sequences discussed above. In summary, the gene sequence was redesigned to eliminate as much as possible the following sequences or sequence features (while avoiding the introduction of unnecessary restriction sites): stretches of G's and C's of 5 or greater; and A+T rich regions (predominantly) that could function as polyadenylation sites or potential RNA destabilization region The sequence of this gene is This coding sequence was shown in Figure 8 (SEQ ID NO:9). expressed in E. coli from the RecA promoter and assayed for EPSPS activity and compared with that from the native CP4 EPSPS gene. The apparent Km for PEP for the native and synthetic genes was 11.8 and 12.7, respectively, indicating that the enzyme expressed from the synthetic gene was unaltered. The N-terminus of the coding sequence was mutagenized to place an SphI site at the ATG to permit the construction of the CTP2-CP4 synthetic fusion for chloroplast import. The following primer was used to accomplish this mutagenesis:

25 GGACGGCTGCATGCACCGTGAAGCATGCTTAAGCTTGGCGT AATCATGG (SEQ ID NO:35).

Expression of Chloroplast Directed CP4 EPSPS

The glyphosate target in plants, the 5-enolpyruvyl-shikimate-3-phosphate synthase (EPSPS) enzyme, is located in the chloroplast. Many chloroplast-localized proteins,

including EPSPS, are expressed from nuclear genes as precursors and are targeted to the chloroplast by a chloroplast transit peptide (CTP) that is removed during the import steps. Examples of other such chloroplast proteins include the small subunit (SSU) of Ribulose-1,5-bisphosphate carboxylase (RUBISCO), Ferredoxin, Ferredoxin oxidoreductase, the Light-harvesting-complex protein I and protein II, and Thioredoxin F. It has been demonstrated in vivo and in vitro that non-chloroplast proteins may be targeted to the chloroplast by use of protein fusions with a CTP and that a CTP sequence is sufficient to target a protein to the chloroplast.

A CTP-CP4 EPSPS fusion was constructed between the Arabidopsis thaliana EPSPS CTP (Klee et al., 1987) and the CP4 EPSPS coding sequences. The Arabidopsis CTP was engineered by site-directed mutagenesis to place a SphI restriction site at the CTP processing site. This mutagenesis replaced the Glu-Lys at this location with Cys-Met. The sequence of this CTP, designated as CTP2 (SEQ ID NO:10), is shown in Figure 9. The N-terminus of the CP4 EPSPS gene was modified to place a SphI site that spans the Met codon. The second codon was converted to one for leucine in this step also. This change had no apparent effect on the in vivo activity of CP4 EPSPS in E. coli as judged by rate of complementation of the aroA allele. This modified N-terminus was then combined with the SacI C-terminus and cloned downstream of the CTP2 sequences. The CTP2-CP4 EPSPS fusion was cloned into pBlueScript KS(+). This vector may be transcribed in vitro using the T7 polymerase and the RNA translated with 35S-Methionine to provide material that may be evaluated for import into chloroplasts isolated from Lactuca sativa using the methods described hereinafter (della-Cioppa et al., 1986, 1987). This template was transcribed in vitro using T7 polymerase and the 35S-methionine-labeled CTP2-CP4 EPSPS material was shown

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to import into chloroplasts with an efficiency comparable to that for the control Petunia EPSPS (control = 35S labeled PreEPSPS [pMON6140; della-Cioppa et al., 1986]).

In another example the Arabidopsis EPSPS CTP,

designated as CTP3, was fused to the CP4 EPSPS through an EcoRI site. The sequence of this CTP3 (SEQ ID NO:12) is shown in Figure 10. An EcoRI site was introduced into the Arabidopsis EPSPS mature region around amino acid 27, replacing the sequence -Arg-Ala-Leu-Leu- with -Arg-Ile-Leu-Leu- in the process. The primer of the following sequence was used to modify the N-terminus of the CP4 EPSPS gene to add an EcoRI site to effect the fusion to the CTP3:

GGAAGACGCCCA<u>GAATTC</u>ACGGTGCAAGCAGCCGG (SEQ ID NO:36) (the *Eco*RI site is underlined).

15 This CTP3-CP4 EPSPS fusion was also cloned into the pBlueScript vector and the T7 expressed fusion was found to also import into chloroplasts with an efficiency comparable to that for the control Petunia EPSPS (pMON6140).

A related series of CTPs, designated as CTP4 (SphI) and CTP5 (EcoRI), based on the Petunia EPSPS CTP and gene were also fused to the SphI- and EcoRI-modified CP4 EPSPS gene sequences. The SphI site was added by site-directed mutagenesis to place this restriction site (and change the amino acid sequence to -Cys-Met-) at the chloroplast processing site. All of the CTP-CP4 EPSPS fusions were shown to import into chloroplasts with approximately equal efficiency. The CTP4 (SEQ ID NO:14) and CTP5 (SEQ ID NO:16) sequences are shown in Figures 11 and 12.

A CTP2-LBAA EPSPS fusion was also constructed following the modification of the N-terminus of the LBAA EPSPS 30 gene by the addition of a SphI site. This fusion was also found to be imported efficiently into chloroplasts.

By similar approaches, the CTP2-CP4 EPSPS and the CTP4-CP4 EPSPS fusion have also been shown to import efficiently into chloroplasts prepared from the leaf sheaths of corn. These results indicate that these CTP-CP4 fusions could also provide useful genes to impart glyphosate tolerance in monocot species.

Those skilled in the art will recognize that various chimeric constructs can be made which utilize the functionality of a particular CTP to import a Class II EPSPS enzyme into the plant cell chloroplast. The chloroplast import of the Class II EPSPS can be determined using the following assay.

Chloroplast Uptake Assay

Intact chloroplasts are isolated from lettuce (Latuca sativa, var. longifolia) by centrifugation in Percoll/ficoll gradients as modified from Bartlett et al (1982). The final pellet of intact chloroplasts is suspended in 0.5 ml of sterile 330 mM sorbitol in 50 mM Hepes-KOH, pH 7.7, assayed for chlorophyll (Arnon, 1949), and adjusted to the final chlorophyll concentration of 4 mg/ml (using sorbitol/Hepes). The yield of intact chloroplasts from a single head of lettuce is 3-6mg chlorophyll.

A typical 300 µl uptake experiment contained 5 mM ATP, 8.3 mM unlabeled methionine, 322 mM sorbitol, 58.3 mM Hepes-KOH (pH 8.0), 50 µl reticulocyte lysate translation products, and intact chloroplasts from L. sativa (200 µg chlorophyll). The uptake mixture is gently rocked at room temperature (in 10 x 75 mm glass tubes) directly in front of a fiber optic illuminator set at maximum light intensity (150 Watt bulb). Aliquot samples of the uptake mix (about 50 µl) are removed at various times and fractionated over 100 µl silicone-oil gradients (in 150 µl polyethylene tubes) by centrifugation at 11,000 X g for 30 seconds. Under these conditions, the intact chloroplasts form a pellet under

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the silicone-oil layer and the incubation medium (containing the reticulocyte lysate) floats on the surface. After centrifugation, the silicone-oil gradients are immediately frozen in dry ice. The chloroplast pellet is then resuspended in 50-100 µl of lysis buffer (10 mM Hepes-KOH pH 7.5, 1 mM PMSF, 1 mM benzamidine, 5 mM e-amino-n-caproic acid, and 30 µg/ml aprotinin) and centrifuged at 15,000 X g for 20 minutes to pellet the thylakoid membranes. The clear supernatant (stromal proteins) from this spin, and an aliquot of the reticulocyte lysate incubation medium from each uptake experiment, are mixed with an equal volume of 2X SDS-PAGE sample buffer for electrophoresis (Laemmli, 1970).

SDS-PAGE is carried out according to Laemmli (1970) in 3-17% (w/v) acrylamide slab gels (60 mm X 1.5 mm) with 3% (w/v) acrylamide stacking gels (5 mm X 1.5 mm). The gel is fixed for 20-30 min in a solution with 40% methanol and 10% acetic acid. Then, the gel is soaked in EN3HANCETM (DuPont) for 20-30 minutes, followed by drying the gel on a gel dryer. The gel is imaged by autoradiography, using an intensifying screen and an overnight exposure to determine whether the CP4 EPSPS is imported into the isolated chloroplasts.

PLANT TRANSFORMATION

Plants which can be made glyphosate tolerant by practice of the present invention include, but are not limited to, soybean, cotton, corn, canola, oil seed rape, flax, sugarbeet, sunflower, potato, tobacco, tomato, wheat, rice, alfalfa and lettuce as well as various tree, nut and vine species.

A double-stranded DNA molecule of the present invention ("chimeric gene") can be inserted into the genome of a plant by any suitable method. Suitable plant transformation

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vectors include those derived from a Ti plasmid of Agrobacterium tumefaciens, as well as those disclosed, e.g., by Herrera-Estrella (1983), Bevan (1984), Klee (1985) and EPO publication 120,516 (Schilperoort et al.). In addition to plant transformation vectors derived from the Ti or root-inducing (Ri) plasmids of Agrobacterium, alternative methods can be used to insert the DNA constructs of this invention into plant cells. Such methods may involve, for example, the use of liposomes, electroporation, chemicals that increase free DNA uptake, free DNA delivery via microprojectile bombardment, and transformation using viruses or pollen.

Class II EPSPS Plant transformation vectors

Class II EPSPS DNA sequences may be engineered into vectors capable of transforming plants by using known techniques. The following description is meant to be illustrative and not to be read in a limiting sense. One of ordinary skill in the art would know that other plasmids, vectors, markers, promoters, etc. would be used with suitable results. The CTP2-CP4 EPSPS fusion was cloned as a BglII-EcoRI fragment into the plant vector pMON979 (described below) to form pMON17110, a map of which is presented in Figure 13. In this vector the CP4 gene is expressed from the enhanced CaMV35S promoter (E35S; Kay et al. 1987). A FMV35S promoter construct (pMON17116) was completed in the following way: The SalI-NotI and the NotI-BglII fragments from pMON979 containing the Spc/AAC(3)-III/oriV and the pBR322/Right Border/NOS 3'/CP4 EPSPS gene segment from pMON17110 were ligated with the XhoI-BglII FMV35S promoter fragment from pMON981. These vectors were introduced into tobacco, cotton and canola.

A series of vectors was also completed in the vector pMON977 in which the CP4 EPSPS gene, the CTP2-CP4 EPSPS fusion, and the CTP3-CP4 fusion were cloned as BglII-SacI fragments to form pMON17124, pMON17119, and pMON17120. respectively. These plasmids were introduced into tobacco. A pMON977 derivative containing the CTP2-LBAA EPSPS gene was also completed (pMON17206) and introduced into tobacco.

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The pMON979 plant transformation/expression vector was derived from pMON886 (described below) by replacing the 10 neomycin phosphotransferase typeII (KAN) gene in pMON886 with the 0.89 kb fragment containing the bacterial gentamicin-3-N-acetyltransferase type III (AAC(3)-III) gene (Hayford et al., 1988). The chimeric P-35S/AA(3)-III/NOS 3' gene encodes gentamicin resistance which permits selection of transformed plant cells. pMON979 also contains a 0.95 kb expression cassette consisting of the enhanced CaMV 35S promoter (Kay et al., 1987), several unique restriction sites, and the NOS 3' end (P-En-CaMV35S/NOS 3'). The rest of the pMON979 DNA segments are exactly the same as in pMON886.

Plasmid pMON886 is made up of the following segments of DNA. The first is a 0.93 kb AvaI to engineered-EcoRV fragment isolated from transposon Tn7 that encodes bacterial spectinomycin/streptomycin resistance (Spc/Str), which is a determinant for selection in E. coli and Agrobacterium 25 tumefaciens. This is joined to the 1.61 kb segment of DNA encoding a chimeric kanamycin resistance which permits selection of transformed plant cells. The chimeric gene (P-35S/KAN/NOS 3') consists of the cauliflower mosaic virus (CaMV) 35S promoter, the neomycin phosphotransferase typeII 30 (KAN) gene, and the 3'-nontranslated region of the nopaline synthase gene (NOS 3') (Fraley et al., 1983). The next segment is

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the 0.75 kb oriV containing the origin of replication from the RK2 plasmid. It is joined to the 3.1 kb SalI to PvuI segment of pBR322 (ori322) which provides the origin of replication for maintenance in E. coli and the bom site for the conjugational transfer into the Agrobacterium tumefaciens cells. The next segment is the 0.36 kb PvuI to BclI from pTiT37 that carries the nopaline-type T-DNA right border (Fraley et al., 1985).

The pMON977 vector is the same as pMON981 except for the presence of the P-En-CaMV35S promoter in place of the FMV35S promoter (see below).

The pMON981 plasmid contains the following DNA segments: the 0.93 kb fragment isolated from transposon Tn7 encoding bacterial spectinomycin/streptomycin resistance [Spc/Str; a determinant for selection in E. coli and Agrobacterium tumefaciens (Fling et al., 1985)]; the chimeric kanamycin resistance gene engineered for plant expression to allow selection of the transformed tissue, consisting of the 0.35 kb cauliflower mosaic virus 35S promoter (P-35S) (Odell et al., 1985), the 0.83 kb neomycin phosphotransferase typeII gene (KAN), and the 0.26 kb 3'-nontranslated region of the nopaline synthase gene (NOS 3') (Fraley et al., 1983); the 0.75 kb origin of replication from the RK2 plasmid (oriV) (Stalker et al., 1981); the 3.1 kb SalI to PvuI segment of pBR322 which provides the origin of replication for maintenance in E. coli (ori-322) and the bom site for the conjugational transfer into the Agrobacterium tumefaciens cells, and the 0.36 kb PvuI to BclI fragment from the pTiT37 plasmid containing the nopaline-type T-DNA right border region (Fraley et al., 1985). The expression cassette consists of the 0.6 kb 35S promoter from the figwort mosaic virus (P-FMV35S) (Gowda et al., 1989) and the 0.7 kb 3' non-translated region of the pea rbcS-E9 gene (E9 3') (Coruzzi et al., 1984, and Morelli et al., 1985). The 0.6 kb SspI fragment

containing the FMV35S promoter (Figure 1) was engineered to place suitable cloning sites downstream of the transcriptional start site. The CTP2-CP4syn gene fusion was introduced into plant expression vectors (including pMON981, to form pMON17131; Figure 14) and transformed into tobacco, canola, potato, tomato, sugarbeet, cotton, lettuce, cucumber, oil seed rape, poplar, and Arabidopsis.

The plant vector containing the Class II EPSPS gene may be mobilized into any suitable Agrobacterium strain for 10 transformation of the desired plant species. The plant vector may be mobilized into an ABI Agrobacterium strain. A suitable ABI strain is the A208 Agrobacterium tumefaciens carrying the disarmed Ti plasmid pTiC58 (pMP90RK) (Koncz and Schell, 1986). The Ti plasmid does not carry the T-DNA phytohormone genes 15 and the strain is therefore unable to cause the crown gall disease. Mating of the plant vector into ABI was done by the triparental conjugation system using the helper plasmid pRK2013 (Ditta et al., 1980). When the plant tissue is incubated with the ABI::plant vector conjugate, the vector is transferred to the plant cells by the 20 vir functions encoded by the disarmed pTiC58 plasmid. The vector opens at the T-DNA right border region, and the entire plant vector sequence may be inserted into the host plant chromosome. The pTiC58 Ti plasmid does not transfer to the plant cells but remains in the Agrobacterium.

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Class II EPSPS free DNA vectors

Class II EPSPS genes may also be introduced into plants through direct delivery methods. A number of direct delivery vectors were completed for the CP4 EPSPS gene. The vector pMON13640, a map of which is presented in Figure 15, is described here. The plasmid vector is based on a pUC plasmid

(Vieira and Messing, 1987) containing, in this case, the nptII gene (kanamycin resistance; KAN) from Tn903 to provide a selectable marker in E. coli. The CTP4-EPSPS gene fusion is expressed from the P-FMV35S promoter and contains the NOS 3' polyadenylation sequence fragment and from a second cassette consisting of the E35S promoter, the CTP4-CP4 gene fusion and the NOS 3' sequences. The scoreable GUS marker gene (Jefferson et al. 1987) is expressed from the mannopine synthase promoter (P-MAS; Velten et al., 1984) and the soybean 7S storage protein gene 3' sequences (Schuler et al., 1982). Similar plasmids could also be made in which CTP-CP4 EPSPS fusions are expressed from the enhanced CaMV35S promoter or other plant promoters. Other vectors could be made that are suitable for free DNA delivery into plants and such are within the skill of the art and contemplated to 15 be within the scope of this disclosure.

PLANT REGENERATION

When expression of the Class II EPSPS gene is 20 achieved in transformed cells (or protoplasts), the cells (or protoplasts) are regenerated into whole plants. Choice of methodology for the regeneration step is not critical, with suitable protocols being available for hosts from Leguminosae (alfalfa, soybean, clover, etc.), Umbelliferae (carrot, celery, parsnip), Cruciferae (cabbage, radish, rapeseed, etc.), Cucurbitaceae (melons and cucumber), Gramineae (wheat, rice, corn, etc.), Solanaceae (potato, tobacco, tomato, peppers), various floral crops as well as various trees such as poplar or apple, nut crops or vine plants such as grapes. See, e.g., Ammirato, 1984; Shimamoto, 1989; Fromm, 1990; Vasil, 1990.

The following examples are provided to better elucidate the practice of the present invention and should not be interpreted in any way to limit the scope of the present invention. Those skilled in the art will recognize that various modifications, truncations, etc. can be made to the methods and genes described herein while not departing from the spirit and scope of the present invention.

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In the examples that follow, EPSPS activity in plants is assayed by the following method. Tissue samples were collected and immediately frozen in liquid nitrogen. One gram of young leaf tissue was frozen in a mortar with liquid nitrogen and ground to a fine powder with a pestle. The powder was then transferred to a second mortar, extraction buffer was added (1 ml /gram), and the sample was ground for an additional 45 seconds. extraction buffer for Canola consists of 100 mM Tris, 1 mM EDTA, 10 % glycerol, 5 mM DTT, 1 mM BAM, 5 mM ascorbate, 1.0 mg/ml BSA, pH 7.5 (4°C). The extraction buffer for tobacco consists of 100 mM Tris, 10 mM EDTA, 35 mM KCl, 20 % glycerol, 5 mM DTT, 1 mM BAM, 5 mM ascorbate, 1.0 mg/ml BSA, pH 7.5 (4°C). The mixture was transferred to a microfuge tube and centrifuged for 5 minutes. The resulting supernatants were desalted on spin G-50 (Pharmacia) columns, previously equilibrated with extraction buffer (without BSA), in 0.25 ml aliquots. The desalted extracts were assayed for EPSP synthase activity by radioactive HPLC assay. Protein concentrations in samples were determined by the BioRad microprotein assay with BSA as the standard.

Protein concentrations were determined using the BioRad Microprotein method. BSA was used to generate a standard curve ranging from 2 - $24~\mu g$. Either 800 μl of standard or diluted sample was mixed with 200 μl of concentrated BioRad

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Bradford reagent. The samples were vortexed and read at A(595) after ~ 5 minutes and compared to the standard curve.

EPSPS enzyme assays contained HEPES (50 mM), shikimate-3-phosphate (2 mM), NH₄ molybdate (0.1 mM) and KF (5 mM), with or without glyphosate (0.5 or 1.0 mM). The assay mix (30 μl) and plant extract (10 μl) were preincubated for 1 minute at 25°C and the reactions were initiated by adding ¹⁴C-PEP (1 mM). The reactions were quenched after 3 minutes with 50 μl of 90% EtOH/0.1M HOAc, pH 4.5. The samples were spun at 6000 rpm and the resulting supernatants were analyzed for ¹⁴C-EPSP production by HPLC. Percent resistant EPSPS is calculated from the EPSPS activities with and without glyphosate.

The percent conversion of 14C labeled PEP to 14C EPSP was determined by HPLC radioassay using a C18 guard column (Brownlee) and an AX100 HPLC column (0.4 X 25 cm, Synchropak) with 0.28 M isocratic potassium phosphate eluant, pH 6.5, at 1 ml/min. Initial velocities were calculated by multiplying fractional turnover per unit time by the initial concentration of the labeled substrate (1 mM). The assay was linear with time up to ~ 3 minutes and 30% turnover to EPSPS. Samples were diluted with 10 mM Tris, 10% glycerol, 10 mM DTT, pH 7.5 (4°C) if necessary to obtain results within the linear range.

In these assays DL-dithiotheitol (DTT), benzamidine (BAM), and bovine serum albumin (BSA, essentially globulin free) were obtained from Sigma. Phosphoenolpyruvate (PEP) was from Boehringer Mannheim and phosphoenol-[1-14C]pyruvate (28 mCi/mmol) was from Amersham.

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EXAMPLE 1

Transformed tobacco plants have been generated with a number of the Class II EPSPS gene vectors containing the CP4 EPSPS DNA sequence as described above with suitable expression of the EPSPS. These transformed plants exhibit glyphosate tolerance imparted by the Class II CP4 EPSPS.

Transformation of tobacco employs the tobacco leaf disc transformation protocol which utilizes healthy leaf tissue about 1 month old. After a 15-20 minutes surface sterilization with 10% Clorox plus a surfactant, the leaves are rinsed 3 times in sterile water. Using a sterile paper punch, leaf discs are punched and placed upside down on MS104 media (MS salts 4.3 g/l, sucrose 30 g/l, B5 vitamins 500X 2 ml/l, NAA 0.1 mg/l, and BA 1.0 mg/l) for a 1 day preculture.

The discs are then inoculated with an overnight culture of a disarmed Agrobacterium ABI strain containing the subject vector that had been diluted 1/5 (ie: about 0.6 OD). The inoculation is done by placing the discs in centrifuge tubes with the culture. After 30 to 60 seconds, the liquid is drained off and the discs were blotted between sterile filter paper. The discs are then placed upside down on MS104 feeder plates with a filter disc to co-culture.

After 2-3 days of co-culture, the discs are transferred, still upside down, to selection plates with MS104 media. After 2-3 weeks, callus tissue formed, and individual clumps are separated from the leaf discs. Shoots are cleanly cut from the callus when they are large enough to be distinguished from stems. The shoots are placed on hormone-free rooting media (MSO: MS salts 4.3 g/l, sucrose 30 g/l, and B5 vitamins 500X 2 ml/l) with selection for the appropriate antibiotic resistance. Root formation occurred in 1-2

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weeks. Any leaf callus assays are preferably done on rooted shoots while still sterile. Rooted shoots are then placed in soil and kept in a high humidity environment (ie: plastic containers or bags). The shoots are hardened off by gradually exposing them to ambient humidity conditions.

Expression of CP4 EPSPS protein in transformed plants

Tobacco cells were transformed with a number of plant vectors containing the native CP4 EPSPS gene, and using different promoters and/or CTP's. Preliminary evidence for expression of the gene was given by the ability of the leaf tissue from antibiotic selected transformed shoots to recallus on glyphosate. In some cases, glyphosate tolerant callus was selected directly following transformation. The level of expression of the CP4 EPSPS was determined by the level of glyphosate tolerant EPSPS activity (assayed in the presence of 0.5 mM glyphosate) or by Western blot analysis using a goat anti-CP4 EPSPS antibody. The Western blots were quantitated by densitometer tracing and comparison to a standard curve established using purified CP4 EPSPS. These data are presented as % soluble leaf protein. The data from a number of transformed plant lines and transformation vectors are presented in Table VI below.

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Table VI Expression of CP4 EPSPS in transformed tobacco tissue

	Vector	Plant #	CP4 EPSPS ** (% leaf protein)
5	pMON17110	25313	0.02
	pMON17110	25329	0.04
	pMON17116	25095	0.02
	pMON17119	25106	0.09
	pMON17119	25762	0.09
10	pMON17119	25767	0.03

** Glyphosate tolerant EPSPS activity was also demonstrated in leaf extracts for these plants.

15 Glyphosate tolerance has also been demonstrated at the whole plant level in transformed tobacco plants. In tobacco, Ro transformants of CTP2-CP4 EPSPS were sprayed at 0.4 lb/acre (0.448 kg/hectare), a rate sufficient to kill control non-transformed tobacco plants corresponding to a rating of 3, 1 and 0 at days 7, 14 and 28, respectively, and were analyzed vegetatively and reproductively (Table VII).

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Table VII Glyphosate tolerance in R_0 tobacco CP4 transformants Spray rate = 0.4 lb/acre (0.448kg/hectare)

	Vector/Plant #		3	Score*	
5		<u>Vegetative</u>			<u>Fertile</u>
		day7	day 14	day 28	
	pMON17110/25313	6	4	2	no
	pMON17110/25329	9	10	10	yes
	pMON17119/25106	9	9	10	yes

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* Plants are evaluated on a numerical scoring system of 0-10 where a vegetative score of 10 represents no damage relative to nonsprayed controls and 0 represents a dead plant. Reproductive scores (Fertile) are determined at 28 days after spraying and are evaluated as to whether or not the plant is fertile.

EXAMPLE 2

20 Canola plants were transformed with the pMON17110, pMON17116, and pMON17131 vectors and a number of plant lines of the transformed canola were obtained which exhibit glyphosate tolerance.

25 Plant Material

Seedlings of *Brassica napus* cv *Westar* were established in 2 inch (~ 5 cm) pots containing Metro Mix 350. They were grown in a growth chamber at 24°C, 16/8 hour photoperiod, light intensity of 400 uEm-2sec-1 (HID lamps). They were fertilized with Peters 20-10-20 General Purpose Special. After 2 1/2 weeks

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they were transplanted to 6 inch (~ 15 cm) pots and grown in a growth chamber at 15/10°C day/night temperature, 16/8 hour photoperiod, light intensity of 800 uEm-2sec-1 (HID lamps). They were fertilized with Peters 15-30-15 Hi-Phos Special.

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Transformation/Selection/Regeneration

Four terminal internodes from plants just prior to bolting or in the process of bolting but before flowering were removed and surfaced sterilized in 70% v/v ethanol for 1 minute, 2% w/v sodium hypochlorite for 20 minutes and rinsed 3 times with sterile deionized water. Stems with leaves attached could be refrigerated in moist plastic bags for up to 72 hours prior to sterilization. Six to seven stem segments were cut into 5mm discs with a Redco Vegetable Slicer 200 maintaining orientation of basal end.

The Agrobacterium was grown overnight on a rotator at 24°C in 2mls of Luria Broth containing 50mg/l kanamycin, 24mg/l chloramphenicol and 100mg/l spectinomycin. A 1:10 dilution was made in MS (Murashige and Skoog) media giving approximately 9x108 cells per ml. This was confirmed with optical density readings at 660 mu. The stem discs (explants) were inoculated with 1.0ml of Agrobacterium and the excess was aspirated from the explants.

The explants were placed basal side down in petri plates containing 1/10X standard MS salts, B5 vitamins, 3% sucrose, 0.8% agar, pH 5.7, 1.0mg/l 6-benzyladenine (BA). The plates were layered with 1.5ml of media containing MS salts, B5 vitamins, 3% sucrose, pH 5.7, 4.0mg/l p-chlorophenoxyacetic acid, 0.005mg/l kinetin and covered with sterile filter paper.

Following a 2 to 3 day co-culture, the explants were transferred to deep dish petri plates containing MS salts, B5

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vitamins, 3% sucrose, 0.8% agar, pH 5.7, 1mg/l BA, 500mg/l carbenicillin, 50mg/l cefotaxime, 200 mg/l kanamycin or 175mg/l gentamicin for selection. Seven explants were placed on each plate. After 3 weeks they were transferred to fresh media, 5 explants per plate. The explants were cultured in a growth room at 25°C, continuous light (Cool White).

Expression Assay

After 3 weeks shoots were excised from the explants. Leaf recallusing assays were initiated to confirm modification of R_o shoots. Three tiny pieces of leaf tissue were placed on recallusing media containing MS salts, B5 vitamins, 3% sucrose, 0.8% agar, pH 5.7, 5.0mg/l BA, 0.5mg/l naphthalene acetic acid (NAA), 500mg/l carbenicillin, 50mg/l cefotaxime and 200mg/l kanamycin or gentamicin or 0.5mM glyphosate. The leaf assays were incubated in a growth room under the same conditions as explant culture. After 3 weeks the leaf recallusing assays were scored for herbicide tolerance (callus or green leaf tissue) or sensitivity (bleaching).

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Transplantation

At the time of excision, the shoot stems were dipped in Rootone® and placed in 2 inch (~ 5 cm) pots containing Metro-Mix 350 and placed in a closed humid environment. They were placed in a growth chamber at 24°C, 16/8 hour photoperiod, 400 u E m-1sec-2(HID lamps) for a hardening-off period of approximately 3 weeks.

The seed harvested from R_o plants is R_1 seed which gives rise to R_1 plants. To evaluate the glyphosate tolerance of an R_o plant, its progeny are evaluated. Because an R_o plant is assumed to be hemizygous at each insert location, selfing results

in maximum genotypic segregation in the R_1 . Because each insert acts as a dominant allele, in the absence of linkage and assuming only one hemizygous insert is required for tolerance expression, one insert would segregate 3:1, two inserts, 15:1, three inserts 63:1, etc. Therefore, relatively few R_1 plants need be grown to find at least one resistant phenotype.

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Seed from an R₀ plant is harvested, threshed, and dried before planting in a glyphosate spray test. Various techniques have been used to grow the plants for R₁ spray evaluations. Tests are conducted in both greenhouses and growth chambers. Two planting systems are used; ~ 10 cm pots or plant trays containing 32 or 36 cells. Soil used for planting is either Metro 350 plus three types of slow release fertilizer or plant Metro 350. Irrigation is either overhead in greenhouses or sub-irrigation in growth chambers. Fertilizer is applied as required in irrigation water. Temperature regimes appropriate for canola were maintained. A sixteen hour photoperiod was maintained. At the onset of flowering, plants are transplanted to ~15 cm pots for seed production.

A spray "batch" consists of several sets of R₁ progenies all sprayed on the same date. Some batches may also include evaluations of other than R₁ plants. Each batch also includes sprayed and unsprayed non-transgenic genotypes representing the genotypes in the particular batch which were putatively transformed. Also included in a batch is one or more non-segregating transformed genotypes previously identified as having some resistance.

Two-six plants from each individual R_o progeny are not sprayed and serve as controls to compare and measure the glyphosate tolerance, as well as to assess any variability not ş

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induced by the glyphosate. When the other plants reach the 2-4 leaf stage, usually 10 to 20 days after planting, glyphosate is applied at rates varying from 0.28 to 1.12 kg/ha, depending on objectives of the study. Low rate technology using low volumes has been adopted. A laboratory track sprayer has been calibrated to deliver a rate equivalent to field conditions.

A scale of 0 to 10 is used to rate the sprayed plants for vegetative resistance. The scale is relative to the unsprayed plants from the same R_o plant. A 0 is death, while a 10 represents no visible difference from the unsprayed plant. A higher number between 0 and 10 represents progressively less damage as compared to the unsprayed plant. Plants are scored at 7, 14, and 28 days after treatment (DAT), or until bolting, and a line is given the average score of the sprayed plants within an R_o plant family.

Six integers are used to qualitatively describe the degree of reproductive damage from glyphosate:

0: No floral bud development

2: Floral buds present, but aborted prior to opening

4: Flowers open, but no anthers, or anthers fail to extrude past petals

6: Sterile anthers

8: Partially sterile anthers

10: Fully fertile flowers

Plants are scored using this scale at or shortly after initiation of flowering, depending on the rate of floral structure development.

Expression of EPSPS in Canola

After the 3 week period, the transformed canola plants were assayed for the presence of glyphosate tolerant EPSPS

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activity (assayed in the presence of glyphosate at 0.5mM). The results are shown in Table VIII.

Table VIII Expression of CP4 EPSPS in transformed Canola plants

	Tubic All Parity		
5	Vector Control	Plant #	% resistant EPSPS activity of leaf extract (at 0.5 mM glyphosate)
			0
	pMON17110	41	47
10	pMON17110	52	28
	pMON17110	71	82
	pMON17110	104	7 5
	pMON17110	172	84
	pMON17110	177	85
	pMON17110	252	29*
15	pMON17110	350	49
	pMON17116	40	25
	pMON17116	99	87
	pMON17116	175	94
-	pMON17116	178	43
20	pMON17116	182	18
	pMON17116	252	69
	pMON17116	298	44*
	pMON17116	332	89
~-	pMON17116	383	97
25	pMON17116	395	52

^{*}assayed in the presence of 1.0 mM glyphosate

 R_1 transformants of canola were then grown in a growth chamber and sprayed with glyphosate at 0.56 kg/ha (kilogram/hectare) and rated vegetatively. These results are shown in Table IXA - IXC. It is to be noted that expression of

glyphosate resistant EPSPS in all tissues is preferred to observe optimal glyphosate tolerance phenotype in these transgenic plants. In the Tables below, only expression results obtained with leaf tissue are described.

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Table IXA Glyphosate tolerance in Class II EPSPS canola R₁ transformants

(pMON17110 = P-E35S; pMON17116 = P-FMV35S; R1 plants; Spray rate = 0.56 kg/ha)

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			Vegeta	ative
		% resistant	Score'	**
	Vector/Plant No.	EPSPS*	day 7	day 14
15	Control Westar	0	5	3
	pMON17110/41	47	6	7
	pMON17110/71	82	6	7
	pMON17110/177	85	9	10
	pMON17116/40	25	9	9
	pMON17116/99	87	9	10
	pMON17116/175	94	9	10
20	pMON17116/178	43	6 .	3
	pMON17116/182	18	9	10
	pMON17116/383	97	9	10

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Table IXB Glyphosate tolerance in Class II EPSPS canola R₁ transformants

(pMON17131 = P-FMV35S; R1 plants; Spray rate = 0.84 kg/ha)

	Vector/Plant No.	Vegetative score** day 14	Reproductive score day 28
	17131/78	10	10
10	17131/102	9	10
	17131/115	9	10
	17131/116	9	10
	17131/157	9	10
	17131/169	10	10
15	17131/255	10	10
15	control Westar	1	0

Table IXC Glyphosate tolerance in Class I EPSPS canola transformants

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(P-E35S; R2 Plants; Spray rate = 0.28 kg/ha)

			Vegetative		
		% resistant	Score**		
	Vector/Plant No.	EPSPS*	day 7	day 14	
	Control Westar	0	4	2	
25	pMON899/715	96	5	6	
	pMON899/744	95	8	8	
	pMON899/794	86	6	4	
	pMON899/818	81	7	8	
	pMON899/885	57	7	6	

^{* %} resistant EPSPS activity in the presence of 0.5 mM glyphosate

^{**} A vegetative score of 10 indicates no damage, a score of 0 is given to a dead plant.

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The data obtained for the Class II EPSPS transformants may be compared to glyphosate tolerant Class I EPSP transformants in which the same promoter is used to express the EPSPS genes and in which the level of glyphosate tolerant EPSPS activity was comparable for the two types of transformants. A comparison of the data of pMON17110 [in Table IXA] and pMON17131 [Table IXB] with that for pMON899 [in Table IXC; the Class I gene in pMON899 is that from A. thaliana (Klee et al., 1987) in which the glycine at position 101 was changed to an alanine] illustrates that the Class II EPSPS is at least as good as that of the Class I EPSPS. An improvement in vegetative tolerance of Class II EPSPS is apparent when one takes into account that the Class II plants were sprayed at twice the rate and were tested as R₁ plants.

EXAMPLE 3

Soybean plants were transformed with the pMON13640 (Figure 15) vector and a number of plant lines of the transformed soybean were obtained which exhibit glyphosate tolerance.

Soybean plants are transformed with pMON13640 by the method of microprojectile injection using particle gun technology as described in Christou et al. (1988). The seed harvested from R_o plants is R_1 seed which gives rise to R_1 plants. To evaluate the glyphosate tolerance of an R_o plant, its progeny are evaluated. Because an R_o plant is assumed to be hemizygous at each insert location, selfing results in maximum genotypic segregation in the R_1 . Because each insert acts as a dominant allele, in the absence of linkage and assuming only one hemizygous insert is required for tolerance expression, one insert

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would segregate 3:1, two inserts, 15:1, three inserts 63:1, etc. Therefore, relatively few R_1 plants need be grown to find at least one resistant phenotype.

Seed from an R_o soybean plant is harvested, and dried before planting in a glyphosate spray test. Seeds are planted into 4 inch (~5cm) square pots containing Metro 350. Twenty seedlings from each Ro plant is considered adequate for testing. Plants are maintained and grown in a greenhouse environment. A 12.5-14 hour photoperiod and temperatures of 30°C day and 24°C night is regulated. Water soluble Peters Pete Lite fertilizer is applied as needed.

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A spray "batch" consists of several sets of R₁ progenies all sprayed on the same date. Some batches may also include evaluations of other than R₁ plants. Each batch also includes sprayed and unsprayed non-transgenic genotypes representing the genotypes in the particular batch which were putatively transformed. Also included in a batch is one or more non-segregating transformed genotypes previously identified as having some resistance.

One to two plants from each individual R_o progeny are not sprayed and serve as controls to compare and measure the glyphosate tolerance, as well as to assess any variability not induced by the glyphosate. When the other plants reach the first trifoliate leaf stage, usually 2-3 weeks after planting, glyphosate is applied at a rate equivalent of 128 oz./acre (8.895kg/ha) of Roundup®. A laboratory track sprayer has been calibrated to deliver a rate equivalent to those conditions.

A vegetative score of 0 to 10 is used. The score is relative to the unsprayed progenies from the same R_{o} plant. A 0 is death, while a 10 represents no visible difference from the

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unsprayed plant. A higher number between 0 and 10 represents progressively less damage as compared to the unsprayed plant. Plants are scored at 7, 14, and 28 days after treatment (DAT). The data from the analysis of one set of transformed and control soybean plants are described on Table X and show that the CP4 EPSPS gene imparts glyphosate tolerance in soybean also.

Table X Glyphosate tolerance in Class I EPSPS sovbean transformants

(P-E35S, P-FMV35S; RO plants; Spray rate = 128 oz./acre)

	Vector/Plant No.		Vegetative score	
		<u>day 7</u>	<u>day 14</u>	day 28
15	13640/40-11	5	6	7
	13640/40-3	9	10	10
	13640/40-7	4	7	7
	control A5403	2	1	0
	controlA5403	1	1	0

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EXAMPLE 4

The CP4 EPSPS gene may be used to select transformed plant material directly on media containing glyphosate. The ability to select and to identify transformed plant material depends, in most cases, on the use of a dominant selectable marker gene to enable the preferential and continued growth of the transformed tissues in the presence of a normally inhibitory substance. Antibiotic resistance and herbicide tolerance genes have been used almost exclusively as such dominant selectable marker genes in the presence of the corresponding antibiotic or herbicide. The nptII/kanamycin selection scheme is

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probably the most frequently used. It has been demonstrated that CP4 EPSPS is also a useful and perhaps superior selectable marker/selection scheme for producing and identifying transformed plants.

A plant transformation vector that may be used in this scheme is pMON17227 (Figure 16). This plasmid resembles many of the other plasmids described infra and is essentially composed of the previously described bacterial replicon system that enables this plasmid to replicate in *E. coli* and to be introduced into and to replicate in *Agrobacterium*, the bacterial selectable marker gene (Spc/Str), and located between the T-DNA right border and left border is the CTP2-CP4 synthetic gene in the FMV35S promoter-E9 3' cassette. This plasmid also has single sites for a number of restriction enzymes, located within the borders and outside of the expression cassette. This makes it possible to easily add other genes and genetic elements to the vector for introduction into plants.

The protocol for direct selection of transformed plants on glyphosate is outlined for tobacco. Explants are prepared for pre-culture as in the standard procedure as described in Example 1: surface sterilization of leaves from 1 month old tobacco plants (15 minutes in 10% clorox + surfactant; 3X dH₂O washes); explants are cut in 0.5 x 0.5 cm squares, removing leaf edges, mid-rib, tip, and petiole end for uniform tissue type; explants are placed in single layer, upside down, on MS104 plates + 2 ml 4COO5K media to moisten surface; pre-culture 1-2 days. Explants are inoculated using overnight culture of Agrobacterium containing the plant transformation plasmid that is adjusted to a titer of 1.2 X 109 bacteria/ml with 4COO5K media. Explants are placed into a centrifuge tube, the Agrobacterium suspension is added and the mixture of bacteria and explants is "Vortexed" on

maximum setting for 25 seconds to ensure even penetration of bacteria. The bacteria are poured off and the explants are blotted between layers of dry sterile filter paper to remove excess bacteria. The blotted explants are placed upside down on MS104 plates + 2ml 4COO5K media + filter disc. Co-culture is 2-3 days. The explants are transferred to MS104 + Carbenicillin 1000 mg/l + cefotaxime 100 mg/l for 3 days (delayed phase). The explants are then transferred to MS104 + glyphosate 0.05 mM + Carbenicillin 1000 mg/l + cefotaxime 100 mg/l for selection phase. At 4-6 weeks shoots are cut from callus and placed on MSO + Carbenicillin 500 mg/l rooting media. Roots form in 3-5 days, at which time leaf pieces can be taken from rooted plates to confirm glyphosate tolerance and that the material is transformed.

The presence of the CP4 EPSPS protein in these transformed tissues has been confirmed by immunoblot analysis of leaf discs. The data from one experiment with pMON17227 is presented in the following: 139 shoots formed on glyphosate from 400 explants inoculated with Agrobacterium ABI/pMON17227; 97 of these were positive on recallusing on glyphosate. These data indicate a transformation rate of 24 per 100 explants, which makes this a highly efficient and time saving transformation procedure for plants. Similar transformation frequencies have been obtained with pMON17131 and direct selection of transformants on glyphosate with the CP4 EPSPS genes has also been shown in other plant species, including Arabidopsis, potato, tomato, cotton, lettuce, and sugarbeet.

The pMON17227 plasmid contains single restriction enzyme recognition cleavage sites (NotI, XhoI, and BstXI) between the CP4 glyphosate selection region and the left border of the vector for the cloning of additional genes and to facilitate the introduction of these genes into plants.

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EXAMPLE 5

The CP4 EPSPS gene has also been introduced into Black Mexican Sweet (BMS) corn cells with expression of the protein and glyphosate resistance detected in callus.

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The backbone for this plasmid was a derivative of the high copy plasmid pUC119 (Viera and Messing, 1987). The 1.3Kb FspI-DraI pUC119 fragment containing the origin of replication was fused to the 1.3Kb Smal-HindIII filled fragment from pKC7 (Rao and Rogers, 1979) which contains the neomycin phosphotransferase type II gene to confer bacterial kanamycin This plasmid was used to construct a monocot expression cassette vector containing the 0.6kb cauliflower mosaic virus (CaMV) 35S RNA promoter with a duplication of the -90 to -300 region (Kay et al., 1987), an 0.8kb fragment containing an intron from a maize gene in the 5' untranslated leader region, followed by a polylinker and the 3' termination sequences from the nopaline synthase (NOS) gene (Fraley et al., 1983). A 1.7Kb fragment containing the 300bp chloroplast transit peptide from the Arabidopsis EPSP synthase fused in frame to the 1.4Kb coding sequence for the bacterial CP4 EPSP synthase was inserted into the monocot expression cassette in the polylinker between the intron and the NOS termination sequence to form the plasmid pMON19653 (Figure 17).

pMON19653 DNA was introduced into Black Mexican Sweet (BMS) cells by co-bombardment with EC9, a plasmid containing a sulfonylurea-resistant form of the maize acetolactate synthase gene. 2.5mg of each plasmid was coated onto tungsten particles and introduced into log-phase BMS cells using a PDS-1000 particle gun essentially as described (Klein et al., 1989).

Transformants are selected on MS medium containing 20ppb chlorsulfuron. After initial selection on chlorsulfuron, the calli can be assayed directly by Western blot. Glyphosate tolerance can be assessed by transferring the calli to medium containing 5mM glyphosate. As shown in Table XI, CP4 EPSPS confers glyphosate tolerance to corn callus.

Table XI. Expression of CP4 in BMS Corn Callus - pMON 19653

10	Line	CP4 expression (% extracted protein)
	284	0.006%
	287	0.036
	290	0.061
15	295	0.073
	299	0.113
	309	0.042
	313	0.003

To measure CP4 EPSPS expression in corn callus, the following procedure was used: BMS callus (3 g wet weight) was dried on filter paper (Whatman#1) under vacuum, reweighed, and extraction buffer (500 μ l/g dry weight; 100 mM Tris, 1 mM EDTA, 10% glycerol) was added. The tissue was homogenized with a 25 Wheaton overhead stirrer for 30 seconds at 2.8 power setting. After centrifugation (3 minutes, Eppendorf microfuge), the supernatant was removed and the protein was quantitated (BioRad Protein Assay). Samples (50 µg/well) were loaded on an SDS PAGE gel (Jule, 3-17%) along with CP4 EPSPS standard (10 ng), 30 electrophoresed, and transferred to nitrocellulose similarly to a previously described method (Padgette, 1987). The nitrocellulose blot was probed with goat anti-CP4 EPSPS IgG, and developed with I-125 Protein G. The radioactive blot was visualized by autoradiography. Results were quantitated by densitometry on an LKB UltraScan XL laser densitomer and are tabulated below in Table X.

Table XII. Glyphosate resistance in BMS Corn Callus using pMON 19653

10	Vector	Experiment	# chlorsulfuron- resistant lines	# cross-resistant to Glyphosate
	19653	253	120	81/ 120 = 67.5 %
	19653	254	80	37/80 = 46%
15	EC9 contro	1 253/254	8	0/8 = 0%

Improvements in the expression of Class I EPSPS could also be achieved by expressing the gene using stronger plant promoters, using better 3' polyadenylation signal sequences, optimizing the sequences around the initiation codon for ribosome loading and translation initiation, or by combination of these or other expression or regulatory sequences or factors. It would also be beneficial to transform the desired plant with a Class I EPSPS gene in conjunction with another glyphosate tolerant EPSPS gene or a gene capable of degrading glyphosate in order to enhance the glyphosate tolerance of the transformed plant.

From the foregoing, it will be seen that this invention is one well adapted to attain all the ends and objects hereinabove set forth together with advantages which are obvious and which are inherent to the invention. It will be understood that certain features and subcombinations are of utility and may be employed without reference to other features and subcombinations. This is contemplated by and is within the scope of the claims.

Since many possible embodiments may be made of the invention without departing from the scope thereof, it is to be understood that all matter herein set forth or shown in the accompanying drawings is to be interpreted as illustrative and not in a limiting sense.

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EXAMPLE 6

The LBAA Class II EPSPS gene has been introduced into plants and also imparts glyphosate tolerance. Data on tobacco transformed with pMON17206 (infra) are presented in Table XIII.

Table XIII - Tobacco Glyphosate Spray Test (pMON17206: E35S - CTP2-LBaa EPSPS: 0.4 lbs/ac)

20	Line	7 Day Rating
	33358	9
	34586	9
	33328	9
	34606	9
	33377	9
	34611	10
25	34607	10
	34601	9
	34589	9
	Samsum	4
	(Control)	

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SEQUENCE LISTING

		
(1) GENE	RAL INF	ORMATION:
(1)	APPLIC	ANT: Barry, Gerard F.
(-/		Kishore, Ganesh M.
		Padgette, Stephen R.
(ii)	TITLE	OF INVENTION: Glyphosate Tolerant
	5-E	nolpyruvylshikimate-3-Phosphate Synthases
(iii)	NUMBER	OF SEQUENCES: 36
(iv)	CORRES	PONDENCE ADDRESS:
	(A) A	DDRESSEE: Dennis R. Hoerner, Jr., Monsanto Co. BB4F
		TREET: 700 Chesterfield Village Parkway
	• •	CITY: St. Louis
•		STATE: Missouri
		COUNTRY: USA
	(F) 2	ZIP: 63198
(v)	COMPUT	TER READABLE FORM:
		MEDIUM TYPE: Floppy disk
		COMPUTER: IBM PC compatible
	(C) C	OPERATING SYSTEM: PC-DOS/MS-DOS
	(D) S	SOFTWARE: PatentIn Release #1.0, Version #1.25
(vi)	CURREN	NT APPLICATION DATA:
• -•	(A) F	APPLICATION NUMBER: US 07/576537
	(B) I	FILING DATE: 31-AUG-1990
	(C) (CLASSIFICATION:
(viii)	ATTOR	NEY/AGENT INFORMATION:
(,		NAME: Hoerner Jr., Dennis R.
		REGISTRATION NUMBER: 30,914
	(C) I	REFERENCE/DOCKET NUMBER: 38-21(10535)
/{w\	TELEC	OMMUNICATION INFORMATION:
(**/		TELEPHONE: (314)537-6099
		TELEFAX: (314)537-6047
	\- ,	()
(2) INE	ORMATIO	ON FOR SEQ ID NO:1:
(i)		NCE CHARACTERISTICS:
		LENGTH: 597 base pairs
		TYPE: nucleic acid
	(C)	STRANDEDNESS: double

(ii) MOLECULE TYPE: DNA (genomic)

(D) TOPOLOGY: linear

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GCCARAAGCT ACAGGAGATC AATGAAGAAT CTTCAATCAA AGTAAACTAC TGTTCCAGCA	240
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(A) LENGTH: 1982 base pairs (B) TYPE: nucleic acid	
(C) STRANDEDNESS: double	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION: 621426	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
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Met Ser His Gly Ala Ser Ser Arg Pro Ala Thr Ala Arg Lys Ser 1 5 10 15	
TOT GGC CTT TOO GGA ACC GTC CGC ATT CCC GGC GAC AAG TCG ATC TCC	154
Ser Gly Leu Ser Gly Thr Val Arg Ile Pro Gly Asp Lys Ser Ile Ser 20 25 30	
CAC CGG TCC TTC ATG TTC GGC GGT CTC GCG AGC GGT GAA ACG CGC ATC	202
His Arg Ser Phe Met Phe Gly Gly Leu Ala Ser Gly Glu Thr Arg Ile	
35 40 45	

-84-

ACC (GGC Gly	CTT Leu 50	CTG Leu	GAA Glu	GGC Gly	GAG Glu	GAC Asp 55	GTC Val	ATC Ile	TAA NBA	ACG Thr	GGC Gly 60	AAG Lys	GCC Ala	ATO	G t	250	
CAG (GCC Ala 65	ATG Met	GJY	GCC Ala	AGG Arg	ATC Ile 70	CGT Arg	AAG Lys	GAA Glu	GGC Gly	GAC Asp 75	ACC Thr	TGG Trp	ATC Ile	AT Il	C e	298	e t
GAT Asp 80	GGC Gly	GTC Val	GGC Gly	AAT Asn	GGC Gly 85	GGC Gly	CTC Leu	CTG Leu	GCG Ala	CCT Pro 90	GAG Glu	GCG Ala	CCG Pro	CTC Leu	As	T P 5	346	
TTC Phe	GGC Gly	TAA neA	GCC Ala	GCC Ala 100	ACG Thr	GGC Gly	TGC Cyb	CGC Arg	CTG Leu 105	ACC Thr	ATG Met	GGC	CTC	GTC Val 110	Gl	G Y	394	
GTC Val	TAC Tyr	GAT Asp	TTC Phe 115	Asp	AGC Ser	ACC Thr	TTC Phe	ATC Ile 120	GGC Gly	GAC Asp	GCC	TCG	Leu 125	Thr	. AA	.G 78	442	
CGC Arg	CCG Pro	ATG Met 130	Gly	CGC Arg	GTG Val	TTG Leu	AAC Asn 135	CCG Pro	CTG Leu	CGC Arg	GAA Glu	ATG Met	Gly	GTG Val	G)	AG Ln	490	
GTG Val	AAA Lys 145	Ser	GAA Glu	GAC Asp	GCT	GAC Asp 150	Arg	CTT Leu	Pro	GTT Val	ACC Thr 155	Lev	G CGC	GGG Gly	Pı	co co	538	
AAG Lys 160	Thr	CCG Pro	ACG Thr	CCG Pro	ATC Ile 165	Thr	TAC Tyr	CGC	GTG Val	Pro	Met	GCC : Ala	TCC Se	C GCA	a G	AG ln 75	586	
GTG Val	AAG Lys	TCC Ser	GCC Ala	GTG Val 180	Lev	CTC Lev	GCC Ala	GGC Gly	CTC Lev 189	AAC 1 ABC	ACC Thi	CC Pr	C GG C G1	C ATO y Ilo 190	e T	CG hr	634	
ACG Thr	GTC Val	ATC	GAC Glu 195	ı Pro	ATC Ile	ATC Met	ACC Thi	Arg 200	J Asj	r CAT P His	r AC	G GA r Gl	A AA u Ly 20	8 We	G C t L	TG eu	682	
CAG Gln	GGC	TT: Pho 210	Gly	GCC Ala	AA(CT n Le	r ACC 2 Thi 215	. Va	C GAG	G ACC	G GA r As	T GC p Al 22	a As	C GG p Gl	С G y V	TG al	730	
CGC	22!	: Il	C CG	C CTO	G GA	A GG(u Gl) 23	y Ar	G GGG G Gl	C AA y Ly	G CT	C AC u Th 23	r Gl	C CA	A GT n Va	C A	TC Le	778	1
GAC Asp 240	Va:	G CC	G GG O Gl	C GA	C CC p Pro 24	o Se	C TC r Se:	G AC	G GC r Al	C TT a Ph 25	e Pr	G CI	rg G1 eu Va	T GC	.a /	SCC Ala 255	826	3
CTG	CT Le	r GT u Va	T CC 1 Pr	G GG o Gl 26	y Se	C GA r As	C GT p Va	C AC 1 Th	C AT	C CT e Le	C AF	C G?	rg Cr	eu Me	rg 1 et 1 70	AAC Asn	874	

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CCC Pro	ACC Thr	CGC A rg	ACC Thr 275	GGC Gly	CTC Leu	ATC Ile	CTG Leu	ACG Thr 280	CTG Leu	CAG Gln	GAA Glu	ATG Met	GGC Gly 285	GCC Ala	GAC Asp	922
ATC Ile	GAA Glu	GTC Val 290	ATC Ile	AAC Asn	CCG Pro	CGC Arg	CTT Leu 295	GCC Ala	GGC	GGC Gly	GAA Glu	GAC Asp 300	GTG Val	GCG Ala	GAC Asp	970
CTG	CGC Arg 305	GTT Val	CGC Arg	TCC Ser	TCC Ser	ACG Thr 310	CTG Leu	AAG Lys	GGC Gly	GTC Val	ACG Thr 315	GTG Val	CCG Pro	GAA Glu	GAC Asp	1018
Arg	GCG Ala	CCT Pro	TCG Ser	ATG Met	ATC Ile 325	GAC Asp	GAA Glu	TAT Tyr	CCG Pro	ATT Ile 330	CTC Leu	GCT Ala	GTC Val	GCC Ala	GCC Ala 335	1066
GCC Ala	TTC Phe	GCG Ala	GAA Glu	GGG Gly 340	GCG Ala	ACC Thr	GTG Val	ATG Met	AAC Asn 345	GGT Gly	CTG Leu	GAA Glu	GAA Glu	CTC Leu 350	CGC Arg	1114
GTC Val	AAG Lys	GAA Glu	AGC Ser 355	GAC Asp	CGC Arg	CTC Leu	TCG Ser	GCC Ala 360	GTC Val	GCC Ala	AAT Asn	GGC Gly	CTC Leu 365	AAG Lys	CTC Leu	1162
AAT Asn	GGC Gly	GTG Val 370	GAT Asp	TGC Cys	GAT Asp	GAG Glu	GGC Gly 375	GAG Glu	ACG Thr	TCG Ser	CTC	GTC Val 380	GTG Val	CGC Arg	GGC Gly	1210
CGC Arg	CCT Pro 385	gac Asp	GGC Gly	AAG Lys	GGG	CTC Leu 390	GGC Gly	AAC Asn	GCC Ala	TCG Ser	GGC Gly 395	GCC Ala	GCC Ala	GTC Val	GCC Ala	1258
ACC Thr 400	CAT His	CTC Leu	GAT Asp	CAC His	CGC Arg 405	ATC Ile	GCC Ala	ATG Met	AGC Ser	TTC Phe 410	CTC Leu	GTC Val	ATG Met	GGC Gly	CTC Leu 415	1306
GTG Val	TCG Ser	GAA Glu	AAC Asn	CCT Pro 420	GTC Val	ACG Thr	GTG Val	GAC Asp	GAT Asp 425	GCC Ala	ACG Thr	ATG Met	ATC Ile	GCC Ala 430	ACG Thr	1354
AGC Ser	TTC Phe	CCG Pro	GAG Glu 435	TTC Phe	ATG Met	GAC Asp	CTG Leu	ATG Met 440	Ala	GGG Gly	CTG Leu	GGC Gly	GCG Ala 445	AAG Lys	ATC Ile	1402
GAA Glu	CTC	TCC Ser 450	GAT Asp	ACG Thr	AAG Lys	GCT Ala	GCC Ala 455	TGA:	rgaco	CTT (CACAI	ATCG	CC A	rcga:	rggtc	1456
CCGC	TGC	GC (CGGC	AAGG	GG A	CCT	CTCG	C GC	CGTA:	rcgc	GGA	GGTC'	TAT (ggct:	TCAT	1516
ATCT	CGAT	rac c	GGC	CTGA	CC T	ATCG	CGCC	A CGO	CCA	AAGC	GCT	CTC	GAT (CGCG	GCCTG?	r 1576
CGCT	TGA?	GA (CGAGO	cccc'	PT G	CGGC	GAT	G TCC	ccc	CAA	TCT	CGAT	CTT (GCCG	GCTC	1636
ACCG	GTC	GT (CTG:	rcgg	ec ci	ATGC	CATC	GCC	BAGG	CGGC	TTC	GAAG	ATC (GCGG:	CATG	1696
CCTC	GGT	CG (CGG	GCGC:	rg G	rcga(GCG	C AG	CGCA	CTT	TGC	GCG	CGT	GAGC	CGGGC	A 1756

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СССТ	CCTG	GA T	GGAC	GCGA'	T AT	CGGC	ACGG	TGG!	TCTG	ccc (GAT	GCGC	CG G	TGAA	GCTCT	,	1816			
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Gly	Leu	Ser	Gly 20	Thr	Val	Arg	Ile	Pro 25	Gly	yab	Lys	Ser	Ile 30	Ser	His					
Arg	Ser	Phe 35	Met	Phe	Gly	Gly	Leu 40	Ala	Ser	Gly	Glu	Thr 45	Arg	Ile	Thr					
Gly	Leu 50	Leu	Glu	Gly	Glu	Asp 55	Val	Ile	Asn	Thr	Gly 60	Lys	Ala	Met	Gln					
Ala 65	Met	Gly	λla	Arg	Ile 70	Arg	Lys	Glu	Gly	Asp 75	Thr	Trp	Ile	Ile	Asp 80					
Gly	Val	Gly	Asn	Gly 85		Leu	Leu	Ala	Pro 90	Glu	Ala	Pro	Leu	Asp 95	Phe					
Gly	Asn	Ala	Ala 100		Gly	Сув	Arg	Leu 105		Жet	Gly	Leu	Val 110		Val					
Tyr	Asp	Phe 115	Авр	Ser	Thr	Phe	Ile 120	Gly	Asp	Ala	Ser	Leu 125	Thr	Lys	Arg					
Pro	Met 130		Arg	Val	Leu	Asn 135	Pro	Leu	Arg	Glu	Met 140		Val	Gln	Val					*
Lys 145		Glu	Авр	Gly	Авр 150		Leu	Pro	Val	Thr 155		Arg	Gly	Pro	Lys 160					1
Thr	Pro	Thr	Pro	Ile 165		Tyr	Arg	Val	Pro 170		Ala	Ser	Ala	Gln 175	Val					
Lye	Ser	Ala	Val 180		Leu	Ala	Gly	Leu 185		Thr	Pro	Gly	11e		Thr					

-87-

Val	Ile	Glu 195	Pro	Ile	Het	Thr	Arg 200	Asp	His	Thr	Glu	L ув 205	Met	Leu	Gln
Gly	Phe 210	Gly	Ala	Aen	Leu	Thr 215	Val	Glu	Thr	Asp	Ala 220	увр	Gly	Val	Arg
Thr 225	Ile	Arg	Leu	Glu	Gly 230	Arg	Gly	Lys	Leu	Thr 235	Gly	Gln	Val	Ile	Авр 240
				245			Thr		250					233	
			260				Thr	265					270		
		275					Thr 280					285			
	290					295	Ala				300				
Arg 305	Val	Arg	Ser	Ser	Thr 310	Leu	Lув	Gly	Val	Thr 315	Val	Pro	Glu	Хвр	Arg 320
Ala	Pro	Ser	Met	Ile 325		Glu	Tyr	Pro	Ile 330	Leu	Ala	Val	Ala	Ala 335	Ala
Phe	Ala	Glu	Gly 340		Thr	Val	Met	Asn 345	Gly	Leu	Glu	Glu	Leu 350	Arg	Val
Lys	Glu	Ser 355		Arg	Leu	Ser	Ala 360	Val	Ala	Asn	Gly	Leu 365	Lys	Leu	Ası
Gly	Val 370		Сув	Авр	Glu	Gly 375	Glu	Thr	Ser	Leu	Val 380	Val	Arg	Gly	Arq
Pro 385	_	Gly	Lys	Gly	Leu 390		Asn	Ala	Ser	Gly 395	Ala	Ala	Val	Ala	400
His	Leu	Asp	His	Arg 405		Ala	Met	Ser	Phe 410	Leu	Val	Met	Gly	Leu 415	Va:
Ser	Glu	Asn	Pro 420		Thr	Val	. Asp	Asp 425	λla	Thr	Met	Ile	Ala 430	Thr	· Se:
Phe	Pro	Glu 435		Met	Авр	Leu	Met 440	Ála	Gly	Leu	Gly	Ala 445	Lys	Ile	Gl
Leu	Ser	Asp	Thr	Lys	Ala	Ala	ı								

(2) INFORMATION FOR SEQ ID NO:4:

450

- (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1673 base pairs

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double

-88-

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 86..1432

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

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TCG GGC GAA ACC CGC ATC ACC (Ser Gly Glu Thr Arg Ile Thr (45			256
ART ACA GGC CGC GCC ATG CAG Asn Thr Gly Arg Ala Met Gln 2			304
GGC GAT GTC TGG ATC ATC AAC Gly Asp Val Trp Ile Ile Asn 75 80		Cys Leu Leu Gln	352
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GAC GCC TCG CTG TCG AAG CGC Asp Ala Ser Leu Ser Lys Arg 125			496
CGC GAA ATG GGC GTT CAG GTG Arg Glu Met Gly Val Gln Val 140			544

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ACC Thr	GAC Asp	AAG Lys 220	GAT Asp	GGC Gly	GTG Val	CGC Arg	CAT His 225	ATC Ile	CGC Arg	ATC Ile	ACC Thr	GGC Gly 230	CAG Gln	GGC Gly	AAG Lys	784
CTT Leu	GTC Val 235	ccc	CAG Gln	ACC Thr	ATC Ile	GAC Asp 240	GTG Val	CCG Pro	GGC	GAT Asp	CCG Pro 245	TCA Ser	TCG Ser	ACC Thr	GCC Ala	832
TTC Phe 250	CCG Pro	CTC Leu	GTT Val	GCC Ala	GCC Ala 255	CTT Leu	CTG Leu	GTG Val	GAA Glu	GGT Gly 260	TCC Ser	GAC Asp	GTC Val	ACC Thr	ATC Ile 265	880
000	***	GTG Val	CTG Leu	ATG Met 270	AAC	CCG Pro	ACC Thr	CGT Arg	ACC Thr 275	GGC Gly	CTC Leu	ATC Ile	CTC Leu	ACC Thr 280	TTG Leu	928
CAG Gln	GAA Glu	ATG Met	GGC Gly 285	GCC	GAT Asp	ATC Ile	GAA Glu	GTG Val 290	CTC Leu	AAT Asn	GCC Ala	CGT Arg	CTT Leu 295	GCA Ala	GGC Gly	976
GGC Gly	GAA Glu	Хвр	GTC Val	GCC Ala	GAT Asp	CTG Leu	CGC Arg	GTC Val	AGG	GCT Ala	TCG Ser	AAG Lys 310	rea	AAG Lys	GGC Gly	1024
GTC Val	Val	Val	CCG	CCG Pro	GAA Glu	CGT Arg 320	GCG Ala	CCG	TCG Ser	ATG Met	ATC Ile 325	Asp	GAA Glu	TAT	CCG Pro	1072
GTC Val 330	Leu	c.CG	ATT Ile	GCC Ala	GCC Ala	TCC	TTC	GCG Ala	GAA Glu	GGC Gly 340	GIU	ACC Thr	GTG Val	ATG Met	GAC Asp 345	1120
	CTC.	GAC Asp	GAA Glu	CTG Leu 350	CGC	GTC	AAG Lys	GAA Glu	TCG Ser 355	yab	CGT	CTG Leu	GCA Ala	GCG Ala 360	GTC Val	1168
GCA Ala	CGC Arg	GGC	CTT Leu 365	GAA Glu	GCC	AAC Asn	GGC	GTC Val	. Asp	т т с с у в	ACC Thr	GAA	GGC Gly 375	GIU	ATG Met	1216

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TCG CTG ACG GTT CGC GGC CG Ser Leu Thr Val Arg Gly Arc 380	C CCC GAC GGC AAG GGA CTG GGC GGC GGC GG Pro Asp Gly Lys Gly Leu Gly Gly Gly 385	1264
ACG GTT GCA ACC CAT CTC GA Thr Val Ala Thr His Leu As 395 40	T CAT CGT ATC GCG ATG AGC TTC CTC GTG P His Arg Ile Ala Met Ser Phe Leu Val 0 405	1312
ATG GGC CTT GCG GCG GAA AA Met Gly Leu Ala Ala Glu Ly 410 415	G CCG GTG ACG GTT GAC GAC AGT AAC ATG 8 Pro Val Thr Val Asp Asp Ser Asn Met 420 425	1360 *
ATC GCC ACG TCC TTC CCC GA Ile Ala Thr Ser Phe Pro Gl 430	A TTC ATG GAC ATG ATG CCG GGA TTG GGC u Phe Met Asp Met Met Pro Gly Leu Gly 435 440	1408
GCA AAG ATC GAG TTG AGC AT Ala Lys Ile Glu Leu Ser Il 445	A CTC TAGTCACTCG ACAGCGAAAA TATTATTTGC e Leu	1462
GAGATTGGGC ATTATTACCG GTTG	GTCTCA GCGGGGGTTT AATGTCCAAT CTTCCATAC	1522
TAACAGCATC AGGAAATATC AAAA	ARGCTT TAGARGGART TGCTAGAGCA GCGACGCCG	1582
CTAAGCTTTC TCAAGACTTC GTTA	AAAACTG TACTGAAATC CCGGGGGTC CGGGGATCA	A 1642
ATGACTTCAT TTCTGAGAAA TTGC	GCCTCGC A	1673
(2) INFORMATION FOR SEQ II	D NO:5:	
(i) SEQUENCE CHARA((A) LENGTH: (B) TYPE: ami (D) TOPOLOGY	449 amino acids ino acid	
(ii) MOLECULE TYPE:	protein	
(xi) SEQUENCE DESCR	IPTION: SEQ ID NO:5:	
Met Ser His Ser Ala Ser P	ro Lys Pro Ala Thr Ala Arg Arg Ser Glu 10 15	
20	rg Ile Pro Gly Asp Lys Ser Ile Ser His 25 30	
35	ly Leu Ala Ser Gly Glu Thr Arg Ile Thr 40 45	•
-	sp Val Ile Asn Thr Gly Arg Ala Met Gln 55 60	3
Ala Met Gly Ala Lys Ile A 65 70	rg Lys Glu Gly Asp Val Trp Ile Ile Asn 75 80	
Gly Val Gly Asn Gly Cys I 85	eu Leu Gln Pro Glu Ala Ala Leu Asp Phe 90 95	

Gly	Asn	Ala	Gly 100	Thr	Gly	Ala	Arg	Leu 105	Thr	Met	Gly	Leu	Val 110	Gly	Thr
Tyr	Asp	Met 115	Lув	Thr	Ser	Phe	11e 120	Gly	Авр	Ala	Ser	Leu 125	Ser	Lys	Arg
Pro	Met 130	Gly	Arg	Val	Leu	Asn 135	Pro	Leu	Arg	Glu	Met 140	Gly	Val	Gln	Val
145					150					133					Lys 160
				165					170						Val
			180					182					1,0		Thr
		195					200					203			Gln
	210					215					220				Arg
225					230					235					Азр 240
				245					250					. 233	
			260					265					2,0		Pro
		275	•				280					200	•		Ile
	290					295	•				300	,			Leu
305					310)				313)				320
				325	i		_		330	,				-	
Phe	Ala	Glu	Gly 340		Thr	Val	Met	Asp 345	Gly	Lev	a yel	Gl:	350	Arç	y Val
Lys	Glu	Sez 355		Arg	, Leu	Ala	360	Val	. Ala	a Arg	g Gly	7 Let 36	ı Glu 5	Ala	a Asn
Gly	Val 370		Cyt	Thr	: Glu	Gly 375	y Glu S	Met	. Sei	. Le	380	va:	l Arq	g Gly	Arç
Pro 385		G13	, Ly	3 Gly	Let 390	. Gly	y Gly	, Gly	Thi	r Val	l Ala 5	Th:	r Hi	s Le	400

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· -92-	
His Arg Ile Ala Met Ser Phe Leu Val Met Gly Leu Ala Ala Glu Lys 405 410 415	
Pro Val Thr Val Asp Asp Ser Asn Met Ile Ala Thr Ser Phe Pro Glu 420 425 430	
Phe Met Asp Met Met Pro Gly Leu Gly Ala Lys Ile Glu Leu Ser Ile 435 440 445	•
Leu	•
(2) INFORMATION FOR SEQ ID NO:6:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 1500 base pairs (B) TYPE: nucleic acid	
(C) STRANDEDNESS: double	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION: 341380	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
GTGATCGCGC CAAAATGTGA CTGTGAAAAA TCC ATG TCC CAT TCT GCA TCC CCG	54
Met Ser His Ser Ala Ser Pro	
	100
ARA CCA GCA ACC GCC CGC CGC TCG GAG GCA CTC ACG GGC GAA ATC CGC	102
Lys Pro Ala Thr Ala Arg Arg Ser Glu Ala Leu Thr Gly Glu Ile Arg 10 15 20	
	150
ATT CCG GGC GAC AAG TCC ATC TCG CAT CGC TCC TTC ATG TTT GGC GGT Ile Pro Gly Asp Lys Ser Ile Ser His Arg Ser Phe Met Phe Gly Gly	
25 30 35	
CTC GCA TCG GGC GAA ACC CGC ATC ACC GGC CTT CTG GAA GGC GAG GAC	198
Tow Ala Ser Gly Glu Thr Arg Ile Thr Gly Leu Leu Glu Gly Gld Asp	
40 45 50 55	
GTC ATC AAT ACA GGC CGC GCC ATG CAG GCC ATG GGC GCG AAA ATC CGT	246
Val Ile Asn Thr Gly Arg Ala Met Gln Ala Met Gly Ala Lys Ite Arg	
60 65 70	
AAA GAG GGC GAT GTC TGG ATC ATC AAC GGC GTC GGC AAT GGC TGC CTG	294
Lys Glu Gly Asp Val Trp Ile Ile Asn Gly Val Gly Asn Gly Cys Led	
75	
TTG CAG CCC GAA GCT GCG CTC GAT TTC GGC AAT GCC GGA ACC GGC GCG	342
Leu Gln Pro Glu Ala Ala Leu Asp Phe Gly Asn Ala Gly Thr Gly Ala	
90 95	

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CGC Arg	CTC Leu 105	ACC Thr	ATG Het	GGC Gly	CTT Leu	GTC Val 110	GGC Gly	ACC Thr	TAT Tyr	GAC Asp	ATG Met 115	AAG Lyb	ACC Thr	TCC Ser	TTT Phe	390
ATC Ile 120	GGC Gly	gac Asp	GCC Ala	TCG Ser	CTG Leu 125	TCG Ser	AÄG Lyb	CGC Arg	CCG Pro	ATG Met 130	GGC Gly	CGC Arg	GTG Val	CTG Leu	AAC Asn 135	438
CCG Pro	TTG Leu	CGC Arg	GAA Glu	ATG Met 140	GGC Gly	GTT Val	CAG Gln	GTG Val	GAA Glu 145	GCA Ala	GCC Ala	GAT Asp	GGC Gly	GAC Asp 150	CGC Arg	486
ATG Met	CCG Pro	CTG Leu	ACG Thr 155	CTG Leu	ATC Ile	GGC	CCG Pro	AAG Lys 160	ACG Thr	GCC Ala	AAT Asn	CCG Pro	ATC Ile 165	ACC Thr	TAT Tyr	534
CGC Arg	GTG Val	CCG Pro 170	ATG Met	GCC Ala	TCC Ser	GCG Ala	CAG Gln 175	GTA Val	AAA Lys	TCC Ser	GCC Ala	GTG Val 180	CTG Leu	CTC Leu	GCC Ala	582
GGT Gly	CTC Leu 185	AAC	ACG Thr	CCG Pro	GCC	GTC Val 190	ACC Thr	ACC Thr	GTC Val	ATC Ile	GAG Glu 195	CCG Pro	GTC Val	ATG Met	ACC Thr	630
CGC Arg 200	GAC Asp	CAC His	ACC Thr	GAA Glu	AAG Lys 205	ATG Met	CTG Leu	CAG Gln	Gly	TTT Phe 210	GGC Gly	GCC Ala	GAC Asp	CTC Leu	ACG Thr 215	678
GTC Val	GAG Glu	ACC Thr	GAC Asp	AAG Lys 220	GAT Asp	GGC Gly	GTG Val	CGC Arg	CAT His 225	ATC Ile	CGC Arg	ATC Ile	ACC Thr	GGC Gly 230	CAG Gln	726
GGC Gly	AAG Lys	CTT Leu	GTC Val 235	GGC Gly	CAG Gln	ACC Thr	ATC Ile	GAC Asp 240	GTG Val	CCG Pro	GGC Gly	GAT Asp	CCG Pro 245	TCA Ser	TCG Ser	774
ACC Thr	GCC Ala	TTC Phe 250	CCG Pro	CTC Leu	GTT Val	GCC Ala	GCC Ala 255	CTT Leu	CTG Leu	GTG Val	GAA Glu	GGT Gly 260	TCC Ser	GAC Asp	GTC Val	822
ACC Thr	ATC Ile 265	CGC Arg	AAC Asn	GTG Val	CTG Leu	ATG Met 270	AAC Asn	CCG Pro	ACC Thr	CGT Arg	ACC Thr 275	GGC Gly	CTC Leu	ATC Ile	CTC Leu	870
ACC Thr 280	TTG Leu	CAG Gln	GAA Glu	ATG Met	GGC Gly 285	GCC Ala	GAT GAT	ATC Ile	GAA Glu	GTG Val 290	CTC Leu	AAT Asn	GCC Ala	CGT Arg	CTT Leu 295	918
GCA Ala	GGC Gly	GGC Gly	GAA Glu	GAC Asp 300	GTC Val	GCC Ala	GAT Asp	CTG Leu	CGC Arg 305	GTC Val	AGG Arg	GCT Ala	TCG Ser	AAG Lys 310	CTC Leu	966
AAG Lyb	GGC Gly	GTC Val	GTC Val 315	GTT Val	CCG Pro	CCG Pro	GAA Glu	CGT Arg 320	GCG Ala	ccg Pro	TCG Ser	ATG Met	ATC Ile 325	GAC Asp	GAA Glu	1014

TAT Tyr	CCG Pro	GTC Val 330	CTG Leu	GCG Ala	ATT Ile	GCC Ala	GCC Ala 335	TCC Ser	TTC Phe	GCG Ala	GAA Glu	GGC Gly 340	GAA Glu	ACC Thr	GT(; L	1062	
ATG Met	GAC Asp 345	GGG Gly	CTC Leu	GAC Asp	GAA Glu	CTG Leu 350	CGC Arg	GTC Val	AAG Lys	GAA Glu	TCG Ser 355	GAT Asp	CGT Arg	CTG Leu	GCI Ala	A 2	1110	i i
GCG Ala 360	GTC Val	GCA Ala	CGC Arg	eja GCC	CTT Leu 365	GAA Glu	GCC Ala	AAC Asn	GGC Gly	GTC Val 370	GAT Asp	TGC Cys	ACC Thr	GAA Glu	GG(G1; 37;	Y	1158	
GAG Glu	ATG Met	TCG Ser	CTG Leu	ACG Thr 380	Val	CGC Arg	GGC	CGC Arg	CCC Pro 385	yab	GGC	AAG Lys	GGA Gly	CTG Leu 390	Gl	Ç Y	1206	
GGC Gly	GGC	ACG Thr	GTT Val 395	Ala	ACC Thr	CAT His	CTC Leu	GAT Asp 400	His	CGT Arg	ATC Ile	GCG Ala	ATG Met 405	Ser	TT Ph	C e	1254	
CTC Leu	GTG Val	ATG Met 410	GGC	CTT Leu	GCG Ala	GCG Ala	GAA Glu 415	Lys	CCG Pro	GTG Val	ACG Thr	GTT Val 420	Asp	GAC Asp	: AG Se	T	1302	
AAC Asn	ATG Met 425	Ile	GCC Ala	ACG Thr	TCC	TTC Phe 430	Pro	GAA Glu	TTC Phe	: ATG : Met	GAC Asp 435	Met	ATG Met	CCG Pro	GG G1	A Y	1350	
	Gly		AAG Lys			Leu					TCAC	TCG	ACAG	CGAF	AA		1400	
TAT	TATT	TGC	GAGA	TTGG	GC A	TTAT	TACC	G GI	TGG	CTC	GCG	GGGG	TTT	AATO	STCC	CAAT	1460	•
CTI	CCAT	'ACG	TAAC	AGCA	TC P	GGAF	ATAI	C A	KAAN	\GCT1	:						1500	
(2)	INF	ORMA	LTION	FOF	r seç) ID	NO: 7	11										
		(i)	(E	1) LE 3) Ti	engti (Pe:	i: 44 amin	TERIS 19 am 10 ac line	nino cid		ds								
	((ii)	MOLE	CULI	E TYI	PE:]	prote	ein				,						
	((xi)	SEQU	JENCI	B DE	CRI	PTIO	N: S	EQ I	D NO	:7:							
	: Sei L	Hi:	s Se		a Se: 5	r Pr	D Ly	B Pr		a Th O	r Al	a Ar	g Ar		r G .5	lu		
Ala	a Le	ı Th	r Gly	-	u Il	e Ar	g Il		o G1 5	у Ав	p Ly	s Se		e Se 10	r H	is		
Ar	g Se	c Ph	e Me ^e 5	t Ph	e Gl	y Gl	y Le		a Se	r Gl	y Gl	u Th	r Ar	g Il	e T	hr		

Gly Leu Leu Glu Gly Glu Asp Val Ile Asn Thr Gly Arg Ala Met Gln Ala Met Gly Ala Lys Ile Arg Lys Glu Gly Asp Val Trp Ile Ile Asn 70 Gly Val Gly Asn Gly Cys Leu Leu Gln Pro Glu Ala Ala Leu Asp Phe Gly Asn Ala Gly Thr Gly Ala Arg Leu Thr Met Gly Leu Val Gly Thr 105 Tyr Asp Met Lys Thr Ser Phe Ile Gly Asp Ala Ser Leu Ser Lys Arg 120 Pro Met Gly Arg Val Leu Asn Pro Leu Arg Glu Met Gly Val Gln Val 135 Glu Ala Ala Asp Gly Asp Arg Met Pro Leu Thr Leu Ile Gly Pro Lys 155 150 Thr Ala Asn Pro Ile Thr Tyr Arg Val Pro Met Ala Ser Ala Gln Val Lys Ser Ala Val Leu Leu Ala Gly Leu Asn Thr Pro Gly Val Thr Thr Val Ile Glu Pro Val Met Thr Arg Asp His Thr Glu Lys Met Leu Gln Gly Phe Gly Ala Asp Leu Thr Val Glu Thr Asp Lys Asp Gly Val Arg His Ile Arg Ile Thr Gly Gln Gly Lys Leu Val Gly Gln Thr Ile Asp Val Pro Gly Asp Pro Ser Ser Thr Ala Phe Pro Leu Val Ala Ala Leu Leu Val Glu Gly Ser Asp Val Thr Ile Arg Asn Val Leu Met Asn Pro Thr Arg Thr Gly Leu Ile Leu Thr Leu Gln Glu Met Gly Ala Asp Ile 280 Glu Val Leu Asn Ala Arg Leu Ala Gly Gly Glu Asp Val Ala Asp Leu Arg Val Arg Ala Ser Lys Leu Lys Gly Val Val Pro Pro Glu Arg 310 Ala Pro Ser Met Ile Asp Glu Tyr Pro Val Leu Ala Ile Ala Ala Ser Phe Ala Glu Gly Glu Thr Val Met Asp Gly Leu Asp Glu Leu Arg Val 345 340

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Lys Glu Ser Asp Arg Leu Ala Ala Val Ala Arg Gly Leu Glu Ala Asn 355 360 365

Gly Val Asp Cys Thr Glu Gly Glu Met Ser Leu Thr Val Arg Gly Arg 370 375 380

Pro Asp Gly Lys Gly Leu Gly Gly Gly Thr Val Ala Thr His Leu Asp 385 390 395 400

His Arg Ile Ala Met Ser Phe Leu Val Met Gly Leu Ala Ala Glu Lys 405 410 415

Pro Val Thr Val Asp Asp Ser Asn Met Ile Ala Thr Ser Phe Pro Glu 420 425 430

Phe Met Asp Met Met Pro Gly Leu Gly Ala Lys Ile Glu Leu Ser Ile 435 440 445

Leu

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 423 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Ser Leu Thr Leu Gln Pro Ile Ala Arg Val Asp Gly Thr Ile Asn Leu 1 5 10 15

Pro Gly Ser Lys Thr Val Ser Asn Arg Ala Leu Leu Leu Ala Ala Leu 20 25 30

Ala His Gly Lys Thr Val Leu Thr Asn Leu Leu Asp Ser Asp Asp Val 35 40 45

Arg His Met Leu Asn Ala Leu Thr Ala Leu Gly Val Ser Tyr Thr Leu 50 55 60

Ser Ala Asp Arg Thr Arg Cys Glu Ile Ile Gly Asn Gly Gly Pro Leu 65 70 75 80

His Ala Glu Gly Ala Leu Glu Leu Phe Leu Gly Asn Ala Gly Thr Ala 85 90 95

Met Arg Pro Leu Ala Ala Ala Leu Cys Leu Gly Ser Asn Asp Ile Val 100 105 110

Leu	ı Thi	Gl ₃		Pro) Arg	Met	Lys 120		Arg	Pro	Ile	Gly 125		Leu	Va
yez) Ala		ı Arç	, Leu	Gly	Gly 135		Lys	Ile	Thr	Tyr 140		Glu	Gln	Gl
Aer 145	_	Pro	Pro	Leu	Arg 150		Gln	Gly	Gly	Phe 155		Gly	Gly	Asn	Va 16
Asp	Val	. Asp	Gly	Ser 165	Val	Ser	Ser	Gln	Phe 170		Thr	Ala	Leu	Leu 175	
Thr	Ala	Pro	180		Pro	Glu	Asp	Thr 185		Ile	Arg	Ile	Lys 190	Gly	Аs
Leu	Val	Ser 195	_	Pro	Tyr	Ile	Asp 200		Thr	Leu	Asn	Leu 205	Met	Lys	Th:
Phe	Gly 210		Glu	Ile	Glu	Asn 215	Gln	His	Tyr	Gln	Gln 220	Phe	Val	Val	Ly
Gly 225		Gln	Ser	Tyr	Gln 230	Ser	Pro	Gly	Thr	Tyr 235	Leu	Val	Glu	Gly	As; 240
Ala	Ser	Ser	λla	Ser 245	Tyr	Phe	Leu	Ala	Ala 250	Ala	Ala	Ile	Lys	Gly 255	G1
Thr	Val	Lys	Val 260	Thr	Gly	Ile	Gly	Arg 265	Asn	Ser	Met	Gln	Gly 270	Asp	Ile
Arg	Phe	Ala 275	_	Val	Leu	Glu	L ув 280	Met	Gly	Ala	Thr	Ile 285	Сув	Trp	Gly
Авр	Asp 290	Tyr	Ile	Ser	Сув	Thr 295	Arg	Gly	Glu	Leu	Asn 300	Ala	Ile	Авр	Met
Авр 305	Met	Asn	His	Ile	Pro 310	Asp	Ala	Ala	Met	Thr 315	Ile	Ala	Thr	Ala	A16
Leu	Phe	Ala	Lys	Gly 325	Thr	Thr	Arg	Leu	Arg 330	Asn	Ile	Tyr	Asn	Trp 335	Arç
Val	Lув	Glu	Thr 340	узр	Arg	Leu	Phe	Ala 345	Met	Ala	Thr	Glu	Leu 350	Arg	Lys
Val	Gly	Ala 355	Glu	Val	Glu	Glu	Gly 360	His	увр	Tyr	Ile	Arg 365	Ile	Thr	Pro
Pro	Glu 370	Lys	Leu	Asn	Phe	Ala 375	Glu	Ile	Ala	Thr	Tyr 380	Asn	Asp	His	Arç
Met 385	Ala	Xet	Сув	Phe	Ser 390	Leu [·]	Val	Ala	Leu	Ser 395	Авр	Thr	Pro	Val	Th:
Ile	Leu	Asp	Pro	Lys	Сув	Thr	Ala		Thr	Phe	Pro	УвЪ	Tyr	Phe	Glu

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Gln Leu Ala Arg Ile Ser Gln 420

(2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1377 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

CCATGGCTCA CGGTGCAAGC AGCCGTCCAG CAACTGCTCG TAAGTCCTCT GGTCTTTCTG 60 GAACCGTCCG TATTCCAGGT GACAAGTCTA TCTCCCACAG GTCCTTCATG TTTGGAGGTC 120 TCGCTAGCGG TGAAACTCGT ATCACCGGTC TTTTGGAAGG TGAAGATGTT ATCAACACTG 180 GTAAGGCTAT GCAAGCTATG GGTGCCAGAA TCCGTAAGGA AGGTGATACT TGGATCATTG 240 ATGGTGTTGG TAACGGTGGA CTCCTTGCTC CTGAGGCTCC TCTCGATTTC GGTAACGCTG 300 CAACTGGTTG CCGTTTGACT ATGGGTCTTG TTGGTGTTTA CGATTTCGAT AGCACTTTCA 360 TTGGTGACGC TTCTCTCACT AAGCGTCCAA TGGGTCGTGT GTTGAACCCA CTTCGCGAAA 420 TGGGTGTGCA GGTGAAGTCT GAAGACGGTG ATCGTCTTCC AGTTACCTTG CGTGGACCAA 480 AGACTCCAAC GCCAATCACC TACAGGGTAC CTATGGCTTC CGCTCAAGTG AAGTCCGCTG 540 TTCTGCTTGC TGGTCTCAAC ACCCCAGGTA TCACCACTGT TATCGAGCCA ATCATGACTC 600 GTGACCACAC TGAAAAGATG CTTCAAGGTT TTGGTGCTAA CCTTACCGTT GAGACTGATG 660 CTGACGGTGT GCGTACCATC CGTCTTGAAG GTCGTGGTAA GCTCACCGGT CAAGTGATTG 720 ATGTTCCAGG TGATCCATCC TCTACTGCTT TCCCATTGGT TGCTGCCTTG CTTGTTCCAG 780 GTTCCGACGT CACCATCCTT AACGTTTTGA TGAACCCAAC CCGTACTGGT CTCATCTTGA 840 CTCTGCAGGA AATGGGTGCC GACATCGAAG TGATCAACCC ACGTCTTGCT GGTGGAGAAG 900 ACGTGGCTGA CTTGCGTGTT CGTTCTTCTA CTTTGAAGGG TGTTACTGTT CCAGAAGACC 960 GTGCTCCTTC TATGATCGAC GAGTATCCAA TTCTCGCTGT TGCAGCTGCA TTCGCTGAAG 1020 GTGCTACCGT TATGAACGGT TTGGAAGAAC TCCGTGTTAA GGAAAGCGAC CGTCTTTCTG 1080 CTGTCGCAAA CGGTCTCAAG CTCAACGGTG TTGATTGCGA TGAAGGTGAG ACTTCTCTCG 1140 TCGTGCGTGG TCGTCCTGAC GGTAAGGGTC TCGGTAACGC TTCTGGAGCA GCTGTCGCTA 1200

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CCCACCTCGA TCACCGTATC GCTATGAGCT TCCTCGTTAT GGGTCTCGTT TCTGAAAACC	1260
CTGTTACTGT TGATGATGCT ACTATGATCG CTACTAGCTT CCCAGAGTTC ATGGATTTGA	1320
TGGCTGGTCT TGGAGCTAAG ATCGAACTCT CCGACACTAA GGCTGCTTGA TGAGCTC	1377
(2) INFORMATION FOR SEQ ID NO:10:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 318 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 87317	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
AGATCTATCG ATAAGCTTGA TGTAATTGGA GGAAGATCAA AATTTTCAAT CCCCATTCTT	60
CGATTGCTTC AATTGAAGTT TCTCCG ATG GCG CAA GTT AGC AGA ATC TGC AAT Met Ala Gln Val Ser Arg Ile Cys Asn 1 5	113
GGT GTG CAG AAC CCA TCT CTT ATC TCC AAT CTC TCG AAA TCC AGT CAA Gly Val Gln Asn Pro Ser Leu Ile Ser Asn Leu Ser Lys Ser Ser Gln 10 15 20 25	161
CGC AAA TCT CCC TTA TCG GTT TCT CTG AAG ACG CAG CAG CAT CCA CGA Arg Lys Ser Pro Leu Ser Val Ser Leu Lys Thr Gln Gln His Pro Arg 30 35 40	209
GCT TAT CCG ATT TCG TCG TCG GGA TTG AAG AAG AGT GGG ATG ACG Ala Tyr Pro Ile Ser Ser Ser Trp Gly Leu Lys Lys Ser Gly Met Thr 45 50 55	257
TTA ATT GGC TCT GAG CTT CGT CCT CTT AAG GTC ATG TCT TCT GTT TCC Leu Ile Gly Ser Glu Leu Arg Pro Leu Lys Val Met Ser Ser Val Ser 60 65 70	305
ACG GCG TGC ATG C Thr Ala Cys Met 75	318
(2) INFORMATION FOR SEQ ID NO:11:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 77 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	•

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(ii) MOLECULE TYPE: protein		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:		
Met Ala Gln Val Ser Arg Ile Cys Asn Gly Val Gln Asn Pro Ser Leu 1 5 10 15		•
Ile Ser Asn Leu Ser Lys Ser Ser Gln Arg Lys Ser Pro Leu Ser Val 20 25 30		4
Ser Leu Lys Thr Gln Gln His Pro Arg Ala Tyr Pro Ile Ser Ser Ser 35 40 45		
Trp Gly Leu Lys Lys Ser Gly Met Thr Leu Ile Gly Ser Glu Leu Arg 50 55 60		
Pro Leu Lys Val Met Ser Ser Val Ser Thr Ala Cys Met 65 70 75		
(2) INFORMATION FOR SEQ ID NO:12:		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 402 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)		
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 87401		٠
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	•	
AGATCTATCG ATAAGCTTGA TGTAATTGGA GGAAGATCAA AATTTTCAAT CCCCATTCTT	60	
CGATTGCTTC AATTGAAGTT TCTCCG ATG GCG CAA GTT AGC AGA ATC TGC AAT Met Ala Gln Val Ser Arg Ile Cys Asn 1 5	113	
GGT GTG CAG AAC CCA TCT CTT ATC TCC AAT CTC TCG AAA TCC AGT CAA Gly Val Gln Asn Pro Ser Leu Ile Ser Asn Leu Ser Lys Ser Ser Gln 10 15 20 25	161	4 .
CGC AAA TCT CCC TTA TCG GTT TCT CTG AAG ACG CAG CAG CAT CCA CGA Arg Lys Ser Pro Leu Ser Val Ser Leu Lys Thr Gln Gln His Pro Arg 30 35 40	209	1
GCT TAT CCG ATT TCG TCG TCG TGG GGA TTG AAG AAG AGT GGG ATG ACG	257	

Ala Tyr Pro Ile Ser Ser Ser Trp Gly Leu Lys Lys Ser Gly Met Thr

50

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TTA Leu	ATT Ile	GGC Gly 60	TCT Ser	GAG Glu	CTT Leu	CGT Arg	CCT Pro 65	CTT Leu	AAG Lys	GTC Val	ATG Met	TCT Ser 70	TCT Ser	GTT Val	TCC Ser	305
ACG Thr	GCG Ala 75	GAG Glu	AAA Lys	GCG Ala	TCG Ser	GAG Glu 80	ATT Ile	GTA Val	CTT Leu	CAA Gln	CCC Pro 85	ATT	AGA Arg	GAA Glu	ATC Ile	353
TCC Ser 90	GGT Gly	CTT Leu	ATT Ile	AAG Lys	TTG Leu 95	CCT Pro	GGC Gly	TCC Ser	AAG Lys	TCT Ser 100	CTA Leu	TCA Ser	AAT Asn	AGA Arg	ATT Ile 105	401
C																402

- (2) INFORMATION FOR SEQ ID NO:13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 105 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met Ala Gln Val Ser Arg Ile Cys Asn Gly Val Gln Asn Pro Ser Leu
1 5 10 15

Ile Ser Asn Leu Ser Lys Ser Ser Gln Arg Lys Ser Pro Leu Ser Val

Ser Leu Lys Thr Gln Gln His Pro Arg Ala Tyr Pro Ile Ser Ser Ser 35 40 45

Trp Gly Leu Lys Lys Ser Gly Met Thr Leu Ile Gly Ser Glu Leu Arg
50 55 60

Pro Leu Lys Val Met Ser Ser Val Ser Thr Ala Glu Lys Ala Ser Glu 65 70 75 80

Ile Val Leu Gln Pro Ile Arg Glu Ile Ser Gly Leu Ile Lys Leu Pro 85 90 95

Gly Ser Lys Ser Leu Ser Asn Arg Ile 100 105

- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 233 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)

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((ix)	(A	TURE () NA () LC	ME/K			232									
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:															
AGAT	GATCTTTCA AGA ATG GCA CAA ATT AAC AAC ATG GCT CAA GGG ATA CAA Met Ala Gln Ile Asn Asn Met Ala Gln Gly Ile Gln 1 5 10														49	
ACC (97
TCA :																145
TCT Ser 1																193
TCC Ser													C			233
(2)	INFO	ORMA:	rion	FOR	SEQ	ID 1	NO:1	5:							٠	
	•	(i) :	(B) LE	NGTH PE:	: 73 amin	ERIS ami: o ac line	no a id								
	(:	ii) l	MOLE	CULE	TYP	E: p	rote	in								
	(:	xi)	SEQU	ence	DES	CRIP	TION	: SE	Q ID	NO:	15:					
Met 1	Ala	Gln	Ile	Asn 5	Asn	Met	Ala	Gln	Gly 10		Gln	Thr	Leu	Asn 15	Pro	
Asn	Ser	Asn	Phe 20		Lys	Pro	Gln	Val 25		Lys	Ser	Ser	Ser 30		Leu	
Val	Phe	Gly 35		Lys	Lys	Leu	Lys 40		Ser	Ala	Asn	Ser 45		Leu	Val	
Leu	Lys 50	_	Asp	Ser	Ile	Phe 55		Gln	Lys	Phe	60 60		Phe	Arg	Ile	
Ser 65	Ala	Ser	Val	Ala	Thr		Cys	Met								

(2) INFORMATION FOR SEQ ID NO:16:

-103-

	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 352 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)															
	(ix)	FE#	TURE	E:		ana										
	(A) NAME/KEY: CDS (B) LOCATION: 49351															
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:															
AGAT	AGATCTGCTA GAAATAATTT TGTTTAACTT TAAGAAGGAG ATATATCC ATG GCA CAA Met Ala Gln 1 ATT AAC AAC ATG GCT CAA GGG ATA CAA ACC CTT AAT CCC AAT TCC AAT 105															
እጥጥ	220	AAC	ATG	GCT	CAA	GGG	ATA	CAA	ACC	CTT	AAT	ccc	AAT	TCC	AAT	105
Ile	Asn 5	Asn	Met	Ala	Gln	Gly 10	Ile	Gln	Thr	Leu	Asn 15	Pro	Asn	Ser	Asn	
TTC	CAT	AAA	ccc	CAA	GTT	CCT	AAA	TCT	TCA	AGT	TTT	CTT	GTT	TTT	GGA	153
Phe	His	Lys	Pro	Gln	Val	Pro	Lys	Ser	Ser	Ser	Phe	Leu	Val	Phe	Gly 35	
20					25					30					-	
TCT	AAA	AAA	CTG	AAA	AAT	TCA	GCA	AAT	TCT	ATG	TTG	GTT	TTG	AAA	AAA	201
Ser	Lys	Lys	Leu	Lys 40	Asn	Ser	Ala	Asn	Ser 45	Met	Leu	Val	Leu	Lys 50	Lys	
GAT	TCA	ATT	TTT	ATG	CAA	AAG	TTT	TGT	TCC	TTT	AGG	ATT	TCA	GCA	TCA	249
yab	Ser	Ile	Phe 55	Met	Gln	Lys	Phe	60 Cys	Ser	Phe	Arg	Ile	Ser 65	Ala	Ser	
GTG	GCT	ACA	GCA	CAG	AAG	CCT	TCT	GAG	ATA	GTG	TTG	CAA	CCC	ATT	AAA	297
Val	Ala	Thr	Ala	Gln	Lys	Pro	Ser	Glu	Ile	Val	Leu	Gln	Pro	Ile	Lys	
		70					75					80				
GAG	ATT	TCA	GGC	ACT	GTT	AAA	TTG	CCT	GGC	TCT	AAA	TCA	TTA	TCT	AAT	345
Glu	Ile	Ser	Gly	Thr	Val	Lys	Leu	Pro	Gly	Ser	Lys	Ser	Leu	Ser	Asn	
	85					90					95					•
AGA	ATT	С														352
Arg	Ile															
100																
						•										
(2)	(2) INFORMATION FOR SEQ ID NO:17:															
		(i) :	SEQU	ENCE	CHA	ract	eris'	TICS	:							•
			()) LE	ngth	: 10	1 am	ino	acid	8						

(2)

- (B) TYPE: amino acid (D) TOPOLOGY: linear

-104-

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Met Ala Gln Ile Asn Asn Met Ala Gln Gly Ile Gln Thr Leu Asn Pro 1 5 10 15

Asn Ser Asn Phe His Lys Pro Gln Val Pro Lys Ser Ser Ser Phe Leu 20 25 30

Val Phe Gly Ser Lys Lys Leu Lys Asn Ser Ala Asn Ser Met Leu Val 35 40 45

Leu Lys Lys Asp Ser Ile Phe Met Gln Lys Phe Cys Ser Phe Arg Ile
50 55 60

Ser Ala Ser Val Ala Thr Ala Gln Lys Pro Ser Glu Ile Val Leu Gln 65 70 75 80

Pro Ile Lys Glu Ile Ser Gly Thr Val Lys Leu Pro Gly Ser Lys Ser 85 90 95

Leu Ser Asn Arg Ile 100

- (2) INFORMATION FOR SEQ ID NO:18:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Xaa His Gly Ala Ser Ser Arg Pro Ala Thr Ala Arg Lys Ser Ser Gly
1 5 10 15

Leu Xaa Gly Thr Val Arg Ile Pro Gly Asp Lys Met 20 25

- (2) INFORMATION FOR SEQ ID NO:19:
 - (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

-105-(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19: Ala Pro Ser Met Ile Asp Glu Tyr Pro Ile Leu Ala Val 5 1 (2) INFORMATION FOR SEQ ID NO:20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: Bingle (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20: Ile Thr Gly Leu Leu Glu Gly Glu Asp Val Ile Asn Thr Gly Lys (2) INFORMATION FOR SEQ ID NO:21: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: 17 ATGATHGAYG ARTAYCC (2) INFORMATION FOR SEQ ID NO:22: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)

17

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

GARGAYGTNA THAACAC

(2) INFORMATION FOR SEQ ID NO:23:

•	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear		•
	(ii) MOLECULE TYPE: DNA (genomic)		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:		
	ARGAYGINA THAATAC	17	
	2) INFORMATION FOR SEQ ID NO:24:		
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 38 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear		
	(ii) MOLECULE TYPE: DNA (genomic)		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	38	
	(2) INFORMATION FOR SEQ ID NO:25:		
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic)		
	(II) NOLECOLE IIFE: DAN (GENOMIC)		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:		
	GGATAGATTA AGGAAGACGC GCATGCTTCA CGGTGCAAGC AGCC	44	•
	(2) INFORMATION FOR SEQ ID NO:26:		.4
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear		
	(ii) MOLECULE TYPE: DNA (genomic)		

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(xi) SEQUENCE DESCRIPTION: SEQ	
GGCTGCCTGA TGAGCTCCAC AATCGCCATC	DATGG 35
(2) INFORMATION FOR SEQ ID NO:27:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pair (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	CB
(ii) MOLECULE TYPE: DNA (genor	nic)
(xi) SEQUENCE DESCRIPTION: SE	
CGTCGCTCGT CGTGCGTGGC CGCCCTGACG	gc 32
(2) INFORMATION FOR SEQ ID NO:28:	
 (i) SEQUENCE CHARACTERISTICS (A) LENGTH: 29 base pai. (B) TYPE: nucleic acid (C) STRANDEDNESS: singl. (D) TOPOLOGY: linear 	rs
<pre>(ii) MOLECULE TYPE: DNA (geno (xi) SEQUENCE DESCRIPTION: SE</pre>	
CGGGCAAGGC CATGCAGGCT ATGGGCGCC	29
(2) INFORMATION FOR SEQ ID NO:29:	
(i) SEQUENCE CHARACTERISTICS (A) LENGTH: 31 base pai (B) TYPE: nucleic acid (C) STRANDEDNESS: singl (D) TOPOLOGY: linear	rs
(ii) MOLECULE TYPE: DNA (geno	omic)
(xi) SEQUENCE DESCRIPTION: SE	
CGGGCTGCCG CCTGACTATG GGCCTCGTCG	G 33
(2) INFORMATION FOR SEQ ID NO:30:	
(i) SEQUENCE CHARACTERISTICS (A) LENGTH: 15 amino ac (B) TYPE: amino acid	s: cids

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	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
	Xaa His Ser Ala Ser Pro Lys Pro Ala Thr Ala Arg Arg Ser Glu 1 5 10 15	
(2)	INFORMATION FOR SEQ ID NO:31:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
-	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
GCGG	GTBGCSG GYTTSGG	17
(2)	INFORMATION FOR SEQ ID NO:32:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: peptide	
	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
	Pro Gly Asp Lys Ser Ile Ser His Arg Ser Phe Met Phe Gly Gly Leu 1 5 10 15	
(2)	INFORMATION FOR SEQ ID NO:33:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 13 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: peptide	

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
	Leu Asp Phe Gly Asn Ala Ala Thr Gly Cys Arg Leu Thr 1 5 10	
(2)	INFORMATION FOR SEQ ID NO:34:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
CGGC	CARTGCC GCCACCGGCG CGCGCC	2
(2)	INFORMATION FOR SEQ ID NO:35:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:	
GGAC	CGGCTGC TTGCACCGTG AAGCATGCTT AAGCTTGGCG TAATCATGG	4
(2)	INFORMATION FOR SEQ ID NO:36:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

GGAAGACGCC CAGAATTCAC GGTGCAAGCA GCCGG

(ii) MOLECULE TYPE: DNA (genomic)

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Claims:

- 1. An isolated DNA sequence encoding an EPSPS enzyme having a K_m for phosphoenolpyruvate (PEP) between 1-150 μ M and a K_i (glyphosate)/ K_m (PEP) ratio between 3-500, which DNA sequence is capable of hybridizing to a DNA probe from a sequence selected from the group consisting of SEQ ID NO:2, SEQ ID NO:4, and SEQ ID NO:6.
- 10 2. A DNA molecule of claim 1 wherein said K_m for phosphoenolpyruvate is between 2-25 μM .
 - 3. A DNA molecule of claim 1 wherein said K_{i}/K_{m} ratio is between 6-250.
 - 4. An isolated DNA sequence encoding a protein which exhibits EPSPS activity wherein said protein is capable of reacting with antibodies raised against a Class II EPSPS enzyme.
- 5. The DNA sequence of Claim 4 wherein said protein is capable of reacting with antibodies raised against a Class II EPSPS enzyme selected from the group consisting of SEQ ID NO:3, SEQ ID NO:5, and SEQ ID NO:7.
- 6. The DNA sequence of Claim 5 wherein said antibodies are raised against a Class II EPSPS enzyme of SEQ ID NO:3.
- 7. A recombinant, double-stranded DNA molecule comprising in sequence:

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- a) a promoter which functions in plant cells to cause the production of an RNA sequence;
- a structural DNA sequence that causes the production of an RNA sequence which encodes a Class II EPSPS enzyme; and
- c) a 3' non-translated region which functions in plant cells to cause the addition of a stretch of polyadenyl nucleotides to the 3' end of the RNA sequence
- where the promoter is heterologous with respect to the structural DNA sequence and adapted to cause sufficient expression of the fusion polypeptide to enhance the glyphosate tolerance of a plant cell transformed with said DNA molecule.
- 15 8. The DNA molecule of Claim 7 in which said structural DNA sequence encodes a fusion polypeptide comprising an amino-terminal chloroplast transit peptide and a Class II EPSPS enzyme.
- 9. The DNA molecule of Claim 8 wherein said structural DNA sequence encoding a Class II EPSPS enzyme is selected from the group consisting of SEQ ID NO:2, SEQ ID NO:4 and SEQ ID NO:6.
- 25 10. The DNA molecule of Claim 9 wherein said sequence is from SEQ ID NO:2.
 - 11. A DNA molecule of Claim 8 in which the promoter is a plant DNA virus promoter.

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- 12. A DNA molecule of Claim 11 in which the promoter is selected from the group consisting of CaMV35S and FMV35S promoters.
- 5 13. A method of producing genetically transformed plants which are tolerant toward glyphosate herbicide, comprising the steps of:
 - a) inserting into the genome of a plant cell a recombinant, double-stranded DNA molecule comprising:
 - i) a promoter which functions in plant cells to cause the production of an RNA sequence,
 - ii) a structural DNA sequence that causes the production of an RNA sequence which encodes a fusion polypeptide comprising an amino terminal chloroplast transit peptide and a Class II EPSPS enzyme.
 - iii) a 3' non-translated DNA sequence which functions in plant cells to cause the addition of a stretch of polyadenyl nucleotides to the 3' end of the RNA sequence

where the promoter is heterologous with respect to the structural DNA sequence and adapted to cause sufficient expression of the fusion polypeptide to enhance the glyphosate tolerance of a plant cell transformed with said gene:

- b) obtaining a transformed plant cell; and
- c) regenerating from the transformed plant cell a genetically transformed plant which has increased tolerance to glyphosate herbicide.

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14. The method of Claim 13 wherein said structural DNA sequence encoding a Class II EPSPS enzyme is selected from the group consisting of SEQ ID NO:2, SEQ ID NO:4, and SEQ ID NO:6.

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- 15. The DNA molecule of Claim 14 wherein said sequence is that as set forth in SEQ ID NO:2.
- 16. A method of Claim 13 in which the promoter is 10 from a plant DNA virus.
 - 17. A method of Claim 16 in which the promoter is selected from the group consisting of CaMV35S and FMV35S promoters.

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- 18. A glyphosate tolerant plant cell comprising a DNA molecule of Claims 8, 9 or 12.
- 19. A glyphosate tolerant plant cell of Claim 18 in20 which the promoter is a plant DNA virus promoter.
 - 20. A glyphosate tolerant plant cell of Claim 19 in which the promoter is selected from the group consisting of CaMV35S and FMV35S promoters.

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21. A glyphosate tolerant plant cell of Claim 18 selected from the group consisting of corn, wheat, rice, soybean, cotton, sugarbeet, oilseed rape, canola, flax, sunflower, potato, tobacco, tomato, alfalfa, poplar, pine, apple and grape.

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- 22. A glyphosate tolerant plant comprising plant cells of Claim 18.
- 23. A glyphosate tolerant plant of Claim 22 in which the promoter is from a DNA plant virus promoter.
 - 24. A glyphosate tolerant plant of Claim 23 in which the promoter is selected from the group consisting of CaMV35S and FMV35S promoters.

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25. A glyphosate tolerant plant of Claim 22 selected from the group consisting of corn, wheat, rice, soybean, cotton, sugarbeet, oilseed rape, canola, flax, sunflower, potato, tobacco, tomato, alfalfa, poplar, pine, apple and grape.

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26. A method for selectively controlling weeds in a field containing a crop having planted crop seeds or plants comprising the steps of:

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a) planting said crop seeds or plants which are glyphosate tolerant as a result of a recombinant double-stranded DNA molecule being inserted into said crop seed or plant, said DNA molecule having:

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ii) a structural DNA sequence that causes the production of an RNA sequence which encodes a polypeptide which comprises an amino terminal chloroplast transit peptide and a Class II EPSPS

i) a promoter which functions in plant cells to cause

the production of an RNA sequence,

enzyme,

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- iii) a 3' non-translated DNA sequence which functions in plant cells to cause the addition of a stretch of polyadenyl nucleotides to the 3' end of the RNA sequence
- where the promoter is heterologous with respect to the structural DNA sequence and adapted to cause sufficient expression of the fusion polypeptide to enhance the glyphosate tolerance of a plant cell transformed with said gene; and
 - b) applying to said crop and weeds in said field a sufficient amount of glyphosate herbicide to control said weeds without significantly affecting said crop.
- 27. The method of Claim 26 wherein said structural DNA sequence encoding a Class II EPSPS enzyme is selected from the sequences as set forth in SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6.
- 28. A method of Claim 27 in which said DNA 20 molecule contains a structural DNA sequence from SEQ ID NO:2.
 - 29. A method of Claim 28 in which said DNA molecule further comprises a promoter selected from the group consisting of the CAMV35SS and FMV35S promoters.
 - 30. A method of Claim 29 in which the crop plant is selected from the group consisting of corn, wheat, rice, soybean, cotton, sugarbeet, oilseed rape, canola, flax, sunflower, potato, tobacco, tomato, alfalfa, poplar, pine, apple and grape.

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6717 6597 6657 6477 6537 6417 GTACGTAGTACCAGTCATTCAAAGTCTTTTTTTTGTAGGTGGCTTCTGAATTTCAATCACC CGTAGAAACTTTCATTAGAACAGTTGTAGCTCGTCGACCGAACACCCCTGGTCTGTTTTT CATGCATCATGGTCAGTAAGTTTCAGAAAAAGACATCCACCGAAGACTTAAAGTTAGTGG CGGTTTTCGATGTCCTCTAGTTACTTAGAAGTTAGTTTCATTTGATGACAAGGTCGT GCATCTTTGAAAGTAATCTTGTCAACATCGAGCAGCTGGCTTGTGGGGACCAGACAAAA **ICCTTCTTAAGAGTCAGGTTTCGGAGTTGTTCCAGTCCCATGTCTCAGAGGTTTGGTAAT** AGGAAGAATTCTCAGTCCAAAGCCTCAACAAGGTCAGGGTACAGAGTCTCCAAACCATTA ACTITATICAAAIIGGIAICGCCAAAACCAAGAAGGAACICCCAICCICAAAGGIIIGIA TGAAATAAGTTTAACCATAGCGGTTTTGGTTCTTCCTTGAGGGTAGGAGTTTCCAAACAT SspI 6538 6598 6658 6478 6418 6358

F 16.

LLL	6837	6897	6954	
AGGAATGGTGCAGAATTGTTAGGCGCACCTACCAAAAGCATCTTTGCCTTTATTGCAAAG +++++	ATAAAGCAGATTCCTCTAGTACAAGTGGGGAACAAATAACGTGGAAAAGAGCTGTCCTG	ACAGCCCACTCACTAATGCGTATGACGAACGCAGTGACGACCACAAAAGAATTCCCTCTA +++++++	SSPI TATAAGAAGGCATTCATTCCCATTTGAAGGATCATCAGATACTAACCAATATTTCTC +++++++	F16. 1(cont.)
6718	6778	6838	8689	

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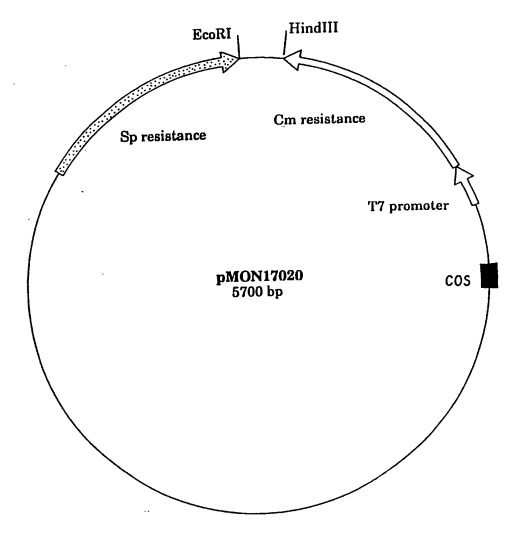


FIG. 2

AAGCCCGCGTTCTCTCCGGCGCTCCGGAGAGCCGTGGATAGATTAAGGAAGACGCC CATGTCGCACGGTGCAAGCGGCCGCAACCGCCCGCAATCCTCTGGCCTTTCCGG M S H G A S S R P A T A R K S S G L S G	Aet) AACCGTCCGCATTCCCGCGACAAGTCGATCTCCCACCGGTCTTCATGTTCGGCGGTCT T V R I P G D K S I S H R S F M F G G L	CGCGAGCGGTGAAACGCGCATCACCGGCCTTCTGGAAGGCGAGGACGTCATCAATACGGG	Ncol Bamhi CAAGGCCATGCAGGCGCCCAGGATCCGTAAGGAAGGCGACACCTGGATCATCGA K A M Q A M G A R I R K E G D T W I I D	CGGCAATGGCGCCTCGTGAGGCGCCCCTCGATTTCGGCAATGCCGC G N G G L L A P E A P L D F G N A A NCOI	GCTGCCGCCTGACCATGGGCCTCGTCGGGTCTACGATTTCGACAGCACCTTCAT	GATGGGCCGCGTTGAACCCGCTGCGCGAAAT M G R V L N P L R E M	TGACCGTCTTCCCGTTACCTTGCGCGGGCCGAA D R L P V T L R G P K	GCCGATGGCCTCCGCACGGTGAAGTCCGCCGT P M A S A Q V K S A V	CATCACGACGGTCATCGAGCCGATCATGACGCG I T T V I E P I M T R
AAGCCCGCGTTCTCTCCGGCGCTCCC CATGTCGCACGGTGCAAGCAGCCGGC	(fMet) AACCGTCCGCATTCCCGGCGACAAG T V R I P G D K S	CGCGAGCGGTGAACGCGCATCACCC	Ncol CAAGGCCATGCAGGCCATGGGCGCC K A M Q A M G A	TGGCGTCGGCAATGGCGGCCTCCTG G V G N G G L L NGOI	CAC	CGCCACCCTCCCTCACAAGCGC G D A S L T K R	GGCCTGCAGGTGAAATCGGAAGACG	GACGCCGACGCCGATCACCTACCGC T P T P I T Y R	S L
61	(f 121	181	241	301	361	421	481	541	601

CGATCATACGGAAAAGATGCTGCAGGGCTTTGGCGCCCAACCTTACCGTCGAGACGGATGC Н G A Sacii ෆ Ø Н Σ × 661

GGACGGCGTGCGCACCATCCGCCTGGAAGGCCGCGGCAAGCTCACCGGCCAAGTCATCGA 721

CGTGCCGGGCGACCCGTCCACGGCCTTCCCGCTGGTTGCGGCCCTGCTTGTTCCGGG 781

> 841

>

GCTGCAGGAAATGGGCGCCGACATCGAAGTCATCAACCCGCGCCTTGCCGGCGGCGAAGA

CGTGGCGGACCTGCGCTTCGCTCCTCCACGCTGAA Ø

CGCGCCTTCGATGATCGAATATCCGATTCTCGCTGTCGCCGCCGCCTTCGCGGAAGG

Ø ⊡ Ω Σ 1021

GGCGACCGTGATGAACGGTCTGGAAGAACTCCGCGTCAAGGAAAGCGACCGCCTCTCGGC CGTCGCCAATGGCCTCAAGCTCAATGGCGTGGATTGCGATGAGGGCGAGACGTCGCTCGT ტ 1081 1141

CGTGCGCGGCCGCCTGACGGCAAGGGGCTCGGCAACGCCTCGGGC > ტ z > 1201

> 1261

1321

1381

961

901

GAAATCCTCGGCAATGGCGGGTTGGCCGATTACGGGACGATCCTCGAGGATATCCGCGGC CGCGACGAGCGGGACATGGGTCGGGCGGACAGTCCTTTGAAGCCCGCCGACGATGCGCAC GATCTTGCCGGGCTCGGTCGGTGCTGTCGGCCCATGCCATCGGCGAGGCGGCTTCG **AAGATCGCGGTCATGCCCTCGGTGCGGCGGCGCTCGTCGAGGCGCAGCGCAGCTTTGCG** GCGCGTGAGCCGGGCACGGTGCTGGATGGACGCGATATCGGCACGGTGGTCTGCCCGGAT GCGCCGGTGAAGCTCTATGTCACCGCGTCACCGGAAGTGCGCGCGAAACGCCGCTATGAC ATCGCCATCGATGGTCCCGCTGCGGCCAAGGGGACGCTCTCGCGCCGTATCGCGGAG GTCTATGGCTTTCATCATCTCGATACGGGCCTGACCTATCGCGCCACGGCCAAAGCGCTG CTCGATCGCGGCCTGTCGCTTGATGACGAGGCGGTTGCGGCCGATGTCGCCCCGCAATCTC 1681 1741 1801 1861 1921 1981 1441 1501 1621 .561

	H M	
84	CAAGO	781
	П	
78	CGACCTCACGGTCGAGACCGACAAGGATGGCGTGCGCCATATCCGCATCACCGGCCAGGG	721
	Ι >	,
72	CGICA	661
	S	,
99	CICCO	601
	Ц	,
9) විධාර	541
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54	CGTGC	481
	Z X	
48	CTATE	421
	A I	
42	TGCGC	361
•	ы	
36	AGAGG	301
	ы	
30	AGGCG	241
	м	
24	TCGCT	181
	ጸ	
18	သဗသ	121
12	GCCAA	61
9	GTAGC	 1

901 GACC T 961 TGCC A 1021 GGGG	VAAALLVEGSDVTIRNVLMNP	
961 TGCC A 1021 GGGC	CGTACCGGCCTCATCCTCACCTTGCAGGAATGGGCGCCGATATCGAAGTGCTCAA	096
1021 GGGC	CGTCTTGCAGGCGCGAAGACGTCGCGCATCTGCGCGCTTCGAAGCTCAA	1020
ני	GTCGTCCTTCCCCCGCAACGTCCATGATCGACGAATATCCGGTCCTGCC	1080
1081 GATT	GCCGCCTCCTTCGCGGAAGCGAAACCGTGATGGACGGGCTCGACGAACTGCGCGT	1140
1141 CAAG	GAATCGGATCGTCTGCCACGCGCCTTGAAGCCAACGCGTCGATTG	1200
1201 CACC	GAAGGCGAGATGTCGCTGATCGCGCCCCCCCGACGCAAGGGACTGGGCGG	1260
1261 CGG(51 CGCCACGGTTGCAACCCATCTCGATCGTATCGCGATGAGCTTCCTCGTGATGGGCCT	1320
1321 TGC(GCGGAAAAGCCGGTGACGTTGACGACAGTAACATGATCGCCACGTCCTTCCCCGA	1380
1381 ATT	ATTCATGGACATGACGGGATTGGGCGCAAAGATCGAGTTGAGCATACTCTAGTCACT	1440
441	CGACAGCGAAATATTTTGCGAGATTGGGCATTATTACCGGTTGGTCTCAGCGGGGT	1500
201	TTAATGTCCAATCTTCCATACGTAACAGCATCAGGAAATATCAAAAAAAGCTTTAGAAGGA	1560
1561 ATT	ATTGCTAGAGCAGGGCGCGCCTAAGCTTTCTCAAGACTTGGTTAAAACTGTACTGAAA TCCCGGGGGGTCCGGGGATCAAATGACTTCATTTCTGAGAAATTGGCCTCGCA	1620 1673
	F16. 4(cont.)	

-	GTGATCGCCCAAAATGTGACTGTGAAAAATCCATGTCCCATTCTGCATCCCCGAAACCA	09
	H S E	
61	CCGCTCGGAGGCACTCACGGGCGAAATCC	120
	RSEALTGEIR	
121	CTCCTTCATGTTTGGCGGTCTCGCATCGG	180
	SFMFGGLASG	
181	CGAGGACGTCATCAATACAGGCCGCGCCA	240
	E D V I N T G R A M	
241	ATCCGTAAAGAGGGCGATGTCTGGATCATCAACGGCGTCGGCAATGGCTGCCTGTTGCAG	300
	G D V W I I N G V G	
301	GCTCGATTTCGGCAATGCCGGAACCGGCG	360
	LDFGNAGTGA	
361	TGACATGAAGACCTCCTTTATCGGCGACG	420
	DMKTSFIGDA	
421	GCTGAACCCGTTGCGCGAAATGGGCGTTC	480
	INPLREMGVO	
481	GCTGACGCTGATCGGCCCGAAGACGGCCA	540
	LTLIGPKTAN	
541	CGCGCAGGTAAATCCGCCGTGCTGCTCG	009
	AQVKSAVLLA	
601	CATCGAGCCGGTCATGACCCGCGACCACA	099
	IEPVMTRDHT	
661	CCTCACGGTCGAGACCGACAAGGATGGCG	720
	LTVETDKDGV	
721	GCTTGTCGGCCAGACCATCGACGTGCCGG	780
781	TGCCGCCCTTCTGGTGGAAGGTTCCGACG	840
	AALLVEGSDV	

GGGCGCCGATATCGAA 900	GADIE	GCGCGTCAGGGCTTCG 960	R V R A S	GATCGACGAATATCCG 1020	товхв	regacegetregaceaa 1080	ОСГОЕ	SCCTTGAAGCCAACGGC 1140	L E A N G	SCCCCGACGCAAGGGA 1200	P D G K G	CGATGAGCTTCCTCGTG 1260	M S F L V	ACATGATCGCCACGTCC 1320	MIATS	TCGAGTTGAGCATACTC 138(пли	ATTACCGGTTGGTCTCA 1440	41 GCGGGGGTTTAATGTCCAATCTTCCATACGTAACAGCATCAGGAAATATCAAAAAAGCTT 1500
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CCT	Н	SCGA	ы	CGGA	더	CGGZ	ഥ	TGGC	Ø	CGC	H	ATC	П	TGA(E	CGG	G	TTT(ATA(
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သဗ္ဗ	i i	SCAG	A G	GTTC	д >	TCCI	S	GATC	DH	GAGP	ᆵ	GCAZ	4	AAGC	×	ATG	Z Z	AAT	ATC
'ACC	H	CTT	H	GTC	>	2225	Æ	ATCG	ຜ	AGGC	ტ	GGTT	>	SGAA	ь	3GAC	Ω	CGAA	ICCA
CCGI	æ	CCGI	ĸ	CGTC	>	TGCC	Ø	GGAI	ы	CGA	臼	CAC	E	3000	Æ	CAT	Σ	CAG	ATG'
GAC	E	MGC	Ø	AGGG	ഗ	CGAT	H	CAA	저	SCAC	H	3000	<u>ග</u>	TTGC	Ø	AATI	ഥ	TCGA	TTTA
ACCC	Д	ICAA	Z	TCA	×	TGGC	K	GCGJ	>	ATT(ပ	900	G	GCC	П	SCG	ы	CAC	GGG
ATGA	N	GTGC	\ \	AAGC	K L	GTCC	V L	CTGC	L R	GICG	O V	CTGG	r G	ATGG	Σ	TTCC	FI P	TAGT	9929
841		901		961		1021		1081		1141		1201		1261		1321		1381	1441

340	294 ISCTRGELNAIDMDMNHIPDAAMTIATAALFAKGTTRLRNIYNWRVK 340:.:::::::::::::::::::::::::::::::	7
303	:. :. :: :. :: :. :: :: :: ::::::::::	7
293	253 AAIKGGTVKVTGIGRNSMQGDIRFADVLEKMGATICWGDDY	7
255	206 MLQGFGANLTVETDADGVRTIRLEGRGKLTGQVIDVPGDPSSTAFPLVAA	0
252		7
205	156 LRGPKTPTPITYRVPMASAQVKSAVLLAGLNTPGITTVIEPIMTRDHTEK	Н
202		7
155	107 MGLVGVYDFDSTFIGDASLTKRPMGRVLNPLREMGVQVK.SEDGDRLPVT	-
152		7
106	59 TGKAMQAMGARIRKEGDTWIIDGVGNGGLLAPEAPLDFGNAATGCRLT	
102	53 MINALTALGVSYTLSADRTRCEIIGNGGPLHAEGALELFLGNAGTAMRPL	
28	• • • • • • • • • • • • • • • • • • •	
52	3 SLTLOPIARVDGTINLPGSKTVSNRALLLAALAHGKTVLTNLLDSDDVRH 52	

F1G. 6(cont.)

300	251 PLVAALLVEGSDVTIRNVLMNPTRTGLILTLOEMGADIEVLNARLAGGED	25
300	գ-	251
250		201
250	Ω-	201
200	: :	151
200	RLPVTLRGPKTPTPITYRVPMASAQVKSAVLLAGLNTPGITTVIEPIMTR	151
150	. .	101
150	TGCRLTMGLVGVYDFDSTFIGDASLTKRPMGRVLNPLREMGVQVKSEDGD	101
100		51
100	LEGEDVINTGKAMQAMGARİRKEGDTWIIDGVGNGGLLAPEAPLDFGNAA	51
50	: : :	
20	MSHGASSRPATARKSSGLSGTVRIPGDKSISHRSFMFGGLASGETRITGL	-

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F16. 7 (cont.)

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Н	CCATGGCTCACGGTGCAAGCAGCCGTCCAGCAACTGCTCGTAAGTCCTCTGGTCTTTCTG	09
61	GAACCGICCGIATICCAGGIGACAAGICIAICICCCACAGGICCIICAIGITIGGAGGIC	120
121	TCGCTAGCGGTGAAACTCGTATCACCGGTCTTTTGGAAGGTGAAGATGTTATCAACACTG	180
181	GTAAGGCTATGCAAGCTATGGGTGCCAGAATCCGTAAGGAAGG	240
241	ATGGTGTTGGTAACGGTGGACTCCTTGCTCCTGAGGCTCCTCTCGATTTCGGTAACGCTG	300
301	CAACTGGTTGCCGTTTGACTATGGGTCTTGTTGGTGTTTACGATTTCGATAGCACTTTCA	360
361	TTGGTGACGCTTCTCTCACTAAGCGTCCAATGGGTCGTGTGTTGAACCCCACTTCGCGAAA	420
421	TGGGTGTGCAGGTGAAGTCTGAAGACGGTGATCGTCTTCCAGTTACCTTGCGTGGACCAA	480
481	AGACTCCAACGCCAATCACCTACAGGGTACCTATGGCTTCCGCTCAAGTGAAGTCCGCTG	540
541	TTCTGCTTGCTGGTCTCAACACCCCCAGGTATCACCACTGTTATCGAGCCAATCATGACTC	009
601	GTGACCACACTGAAAAGATGCTTCAAGGTTTTGGTGCTAACCTTACCGTTGAGACTGATG	660
661	CTGACGGTGTGCGTACCATCCGTCTTGAAGGTCGTGGTAAGCTCACCGGTCAAGTGATTG	720
721	ATGTTCCAGGTGATCCATCCTCTACTGCTTTCCCATTGGTTGCTGCCTTGCTTG	780
781	GTTCCGACGTCACCATCCTTAACGTTTTGATGAACCCAACCCGTACTGGTCTTCATCTTGA	840

F 6.

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137	TGGCTGGTCTTGGAGCTAAGATCGAACTCTCCGACACTAAGGCTGCTTGATGAGCTC	321
132	CTGTTACTGTTGATGCTACTATGATCGCTACTAGCTTCCCAGAGTTCATGGATTTGA	261
126	CCCACCTCGATCACCGTATCGCTATGAGCTTCCTCGTTATGGGTCTCGTTTCTGAAAACC	201
120	TCGTGCGTGGTCGTCTGACGGTAAGGGTCTCGGTAACGCTTCTGGAGCAGCTGTCGCTA	141
114	CTGTCGCAAACGGTCTCAAGCTCAACGGTGTTGATTGCGATGAAGGTGAGACTTCTCTCG	081
108	GTGCTACCGTTATGAACGGTTTGGAAGAACTCCGTGTTAAGGAAAGCGACCGTCTTTCTG	021
102(GTGCTCCTTCTATGATCGACGAGTATCCAATTCTCGCTGTTGCAGCTGCATTCGCTGAAG	961
096	ACGTGGCTGACTTGCTTCGTTCTTCTACTTTGAAGGGTGTTACTGTTCCAGAAGACC	901
900	CTCTGCAGGAAATGGGTGCCGACATCGAAGTGATCAACCCACGTCTTGCTGGTGGAGAAG	841

F16.8(cont.)

F16.9

240 MetAlaGlnValSerArgIleCysAsnGlyValGln AsnProSerLeuIleSerAsnLeuSerLysSerSerGlnArgLysSerProLeuSerVal TCTTGGGTAGAGATAGAGGTTAGAGCTTTAGGTCAGTTGCGTTTAGAGGGAATAGCC AAAGAGACTTCTGCGTCGTAGGTGCTCGAATAGGCTAAAGCAGCAGCACCCCTAACT GCTAACGAAGTTAACTTCAAAGAGGCTACCGCGTTCAATCGTCTTAGACGTTACCACACG **AGAACCCATCTCTTATCTCCAATCTCGAAATCCAGTCAACGCAAATCTCCCTTATCGG** TTTCTCTGAAGACGCAGCATCCACGAGCTTATCCGATTTCGTCGTCGTGGGGATTGA TCTAGATAGCTATTCGAACTACATTAACCTCCTTCTAGTTTTAAAAGTTAGGGGTAAGAA CGATTGCTTCAATTGAAGTTTCTCCGATGGCGCAAGTTAGCAGAATCTGCAATGGTGTGC AGATCTATCGATAAGCTTGATGTAATTGGAGGAAGATCAAAATTTTCAATCCCCATTCTT 61 121

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SerLeuLysThrGlnGlnHisProArgAlaTyrProIleSerSerTrpGlyLeuLys

AGAAGAGTGGGATGACGTTAATTGGCTCTGAGCTTCGTCCTCTTAAGGTCATGTCTTCTG 241

TCTTCTCACCCTACTGCAATTAACCGAGACTCGAAGCAGGAGAATTCCAGTACAGAAGAC

LysSerGlyMetThrLeuIleGlySerGluLeuArgProLeuLysValMetSerSerVal

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AAAGGTGCCGCACGTACG

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TTTCCACGCCGTGCATGC

SerThrAlaCysMet

F16. 9(cont.)

240 180 120 09 MetAlaGlnValSerArgIleCysAsnGlyValGln AsnProSerLeuIleSerAsnLeuSerLysSerSerGlnArgLysSerProLeuSerVal TCTTGGGTAGAGATAGAGGTTAGAGCTTTAGGTCAGTTGCGTTTAGAGGGAATAGCC GCTAACGAAGTTAACTTCAAAGAGGCTACCGCGTTCAATCGTCTTAGACGTTACCACACG AGAACCCATCTCTTATCTCCAATCTCGAAATCCAGTCAACGCAAATCTCCCTTATCGG TITCICIGAAGACGCAGCATCCACGAGCITATCCGATTTCGTCGTCGTGGGGATTGA **AAAGAGACTTCTGCGTCGTAGGTGCTCGAATAGGCTAAAGCAGCAGCACCCCTAACT** CGATTGCTTCAATTGAAGTTTCTCCGATGGCGCAAGTTAGCAGAATCTGCAATGGTGTGC **AGATCTATCGATAAGCTTGATGTAATTGGAGGAAGATCAAAATTTTCAATCCCCATTCTT** TCTAGATAGCTATTCGAACTACATTAACCTCCTTCTAGTTTTAAAAGTTAGGGGTAAGAA 61 121 181

F16. 10(cont.

360 300 SerLeuLysThrGlnGlnHisProArgAlaTyrProIleSerSerSerTrpGlyLeuLys SerThrAlaGluLysAlaSerGluIleValLeuGlnProIleArgGluIleSerGlyLeu LysSerGlyMetThrLeuIleGlySerGluLeuArgProLeuLysValMetSerSerVal TCTTCTCACCCTACTGCAATTAACCGAGACTCGAAGCAGGAGAATTCCAGTACAGAAGAC **AAAGGTGCCGCCTCTTTCGCAGCCTCTAACATGAAGTTGGGTAATCTCTTTAGAGGCCAG AGAAGAGTGGGATGACGTTAATTGGCTCTGAGCTTCGTCCTCTTAAGGTCATGTCTTCTG** TTTCCACGGCGGAGAAAGCGTCGGAGATTGTACTTCAACCCATTAGAGAAATCTCCGGTC **AATAATTCAACGGACCGAGGTTCAGAGATAGTTTATCTTAAG** TTATTAAGTTGCCTGGCTCCAAGTCTCTATCAAATAGAATTC IleLysLeuProGlySerLysSerLeuSerAsnArgIle 国との民 301 361 241

180 233 120 9 LysLysLeuLysAsnSerAlaAsnSerMetLeuValLeuLysLysAspSerIlePheMet MetAlaGlnIleAsnAsnMetAlaGlnGlyIleGlnThrLeuAsnPro AsnSerAsnPheHisLysProGlnValProLysSerSerSerPheLeuValPheGlySer **ATTTTTTGACTTTTAAGTCGTTTAAGATACAACCAAAACTTTTTTCTAAGTTAAAATA** GTTAAGGTTAAAGGTATTTGGGGTTCAAGGATTTAGAAGTTCAAAAGAACAAAAACCTAG <u>Taaaaaactgaaaattcagcaaattctatgttggtttttgaaaaaagattcaatttttt</u> **AGATCTTTCAAGAATGGCACAAATTAACAACATGGCTCAAGGGATACAAACCCTTAATCC** TCTAGAAAGTTCTTACCGTGTTTAATTGTTGTACCGAGTTCCCTATGTTTGGGAATTAGG CAATICCAATITCCATAAACCCCAAGIICCTAAAICIICAAGIITICIIGIITITGGAIC CGTTTTCAAAACAAGGAAATCCTAAAGTCGTAGTCACCGATGTCGGACGTACG GCAAAAGTTTTGTTCCTTTAGGATTTCAGCATCAGTGGCTACAGCCTGCATGC GlnLysPheCysSerPheArgIleSerAlaSerValAlaThrAlaCysMet 181 61 121

> r 7		⊃ -
0.00	TCTATGTTGGTTTTGAAAAAAATTTTTTTTTTTTTTTTT	,
t	ValProLysSerSerPheLeuValPheGlySerLysLysLeuLysAsnSerAlaAsn	
001	CAAGGATTT	7
0	_	101
ı	AsnAsnMetAlaGlnGlyIleGlnThrLeuAsnProAsnSerAsnPheHisLysProGln	
041	TIGITGIAC	Þ
. 001	AACAACATGGCTCAAGGGATACAAACCCTTAATCCCAATTCCAATTTCCATAAACCCCAA	Ü
1	MetAlaGlnIle	
0	TCTAGACGATCTTTATTAAAACAAATTGAAATTCTTCCTCTATATAGGTACCGTGTTTAA	-
Ċ,	AGATCTGCTAGAAATAATTTTGTTTAACTTTTAAGAAGGAGATATATCCATGGCACAAATT	•

- 16. 12

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300 SerMetLeuValLeuLysLysAspSerIlePheMetGlnLysPheCysSerPheArglle AGTCGTAGTCACCGATGTCGTGTCTTCGGAAGACTCTATCACAACGTTGGGTAATTTCTC TCAGCATCAGTGGCTACAGCACAGAAGCCTTCTGAGATAGTGTTGCAACCCCATTAAAGAG 241

SerAlaSerValAlaThrAlaGlnLysProSerGluIleValLeuGlnProIleLysGlu

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TAAAGTCCGTGACAATTTAACGGACCGAGATTTAGTAATAGATTATCTTAAG IleSerGlyThrValLysLeuProGlySerLysSerLeuSerAsnArgIle

ATTICAGGCACTGTTAAATTGCCTGGCTCTAAATCATTATCTAATAGAATTC

301

F16. 12 (cont.

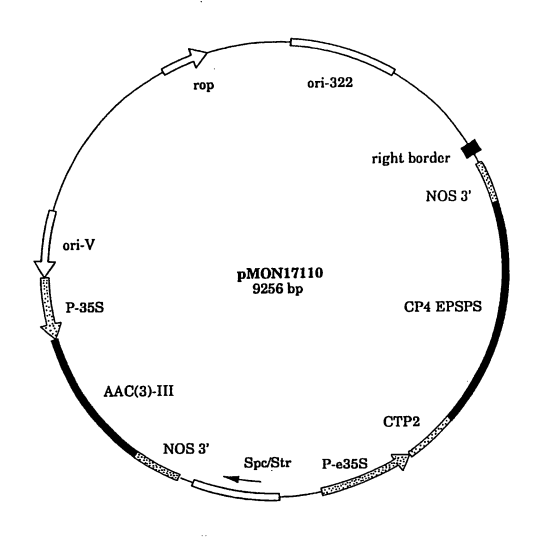


FIG. 13

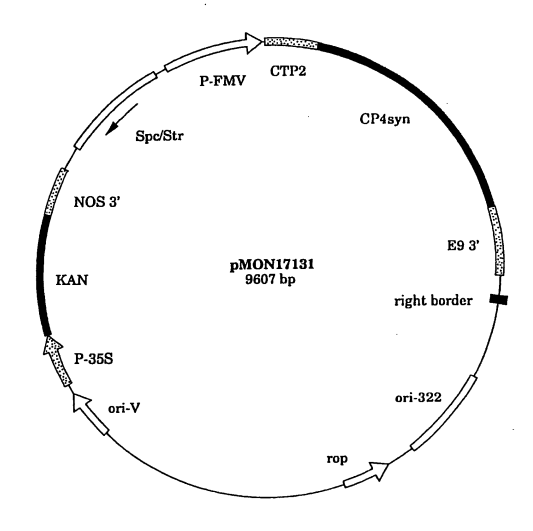
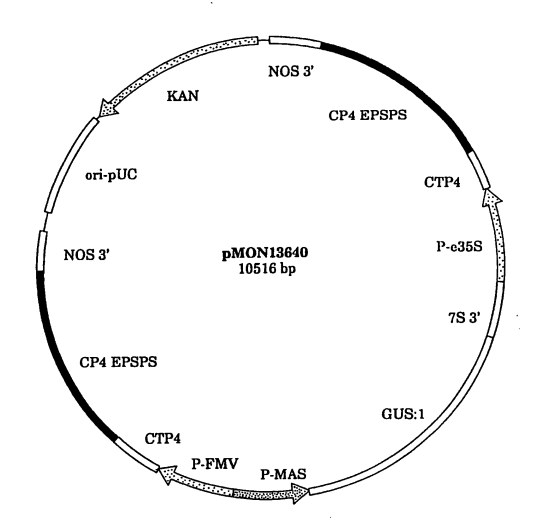


FIG. 14



F I G. 15

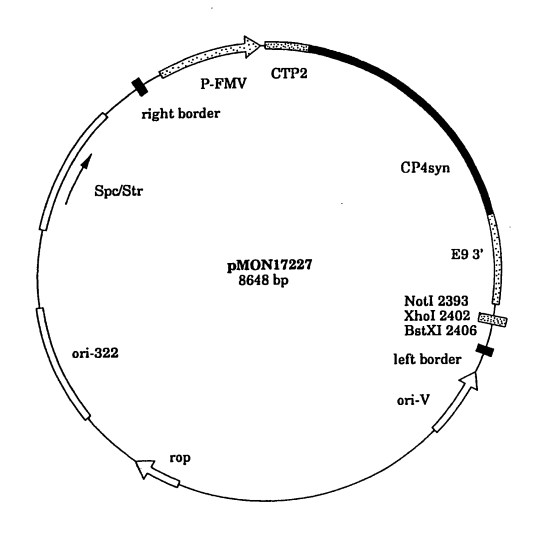


FIG. 16

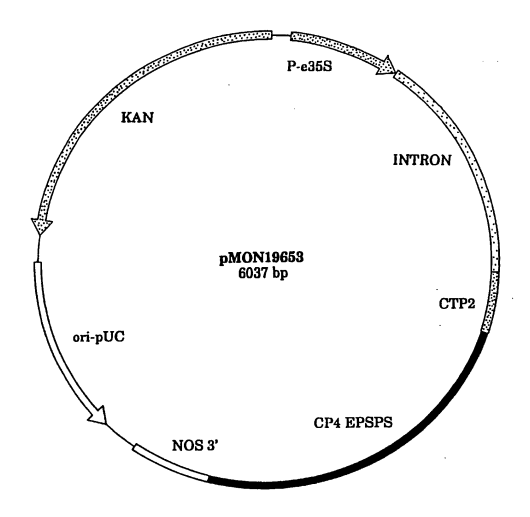


FIG. 17

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 91/06148

		CT MATTER (if several classification sy				
	to International Patent . 5 C12N15/5	Classification (IPC) or to both National Cl 4; C12N15/82;	essification and IPC C12N5/10;	A01H5/00		
II. FIELDS	SEARCHED					
		Minimum Docume	ntation Searched?			
Classificat	tion System		Classification Symbols	·		
Int.Cl	. 5	C12N ; A01H				
		Documentation Searched other to the Extent that such Documents :	than Minimum Documentation ure included in the Fields Searched	1		
III. DOCU		D TO BE RELEVANT ⁹				
Category *	Citation of Do	current, 11 with indication, where appropris	ite, of the relevant passages 12	Relevant to Claim No.13		
A	EP,A,O a	218 571 (MONSANTO) 15 A	pril 1987	1-30		
A	EP,A,O a	293 358 (MONSANTO) 30 N whole document	ovember 1988	1-30		
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			-/			
"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention filing date and not in conflict with the application but cited to understand the principle or theory underlying the invention filing date. "E" earlier document but published on or after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention cannot be committed invention cannot be considered novel or cannot be considered to involve an inventive step when the document referring to an oral disclosure, use, exhibition or other means "P" document published after the international filing date but later than the priority date claimed." "A" document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention cannot be considered novel or cannot be considered to involve an inventive step when the document is combined with one or more other such document, such combination being obvious to a person skilled in the art. "A" document member of the same patent family						
IV. CERTI	FICATION					
Date of the Actual Completion of the International Search 11 DECEMBER 1991			Date of Mailing of this International Search Report 7. 01, 92			
Internations	Searching Authority EUROPE	IN PATENT OFFICE	Signature of Authorized Officer MADDOX A.D.			

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see the whole document		see the whole document	
			
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Form PCT/ISA/210 (extra short) (January 1985)

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO. US 51938

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information. 11/12/91

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